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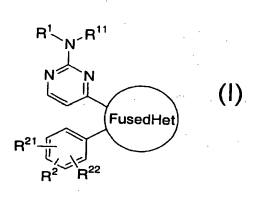
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(54) Title: (PYRIMIDYL)(PHENYL)SUBSTITUTED FUSED HETEROARYL P38 INHIBITING AND PKG KINASE INHIBITING COMPOUNDS



(57) Abstract: Compounds of formula (I) and pharmaceutically acceptable salts thereof are useful in the treatment of cytokine mediated diseases such as arthritis and in the treatment and/or prevention of protozoal diseases such as coccidiosis.

TITLE OF THE INVENTION

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(PYRIMIDYL)(PHENYL)SUBSTITUTED FUSED HETEROARYL P38 INHIBITING AND PKG KINASE INHIBITING COMPOUNDS

5 BACKGROUND OF THE INVENTION

The present invention relates to (pyrimidyl)(phenyl)substituted fused heteroaryl compounds which have cytokine inhibitory activity. The present invention also relates to (pyrimidyl)(phenyl)substituted fused heteroaryl compounds which have cGMP dependent protein kinase ("PKG") inhibitory activity.

Cytokine mediated diseases and cytokine inhibition, suppression and antagonism are used in the context of diseases or conditions in which excessive or unregulated production or activity of one or more cytokines occurs. Examples of cytokines which are effected typically include Interleukin-1 (IL-1), Interleukin-6 (IL-6), Interleukin-8 (IL-8) and Tumor Necrosis Factor (TNF).

Interleukin-1 (IL-1) and Tumor Necrosis Factor (TNF) are produced by a variety of cells that are involved in immunoregulation and other physiological conditions.

There are many disease states in which IL-1 is implicated. Examples are rheumatoid arthritis, osteoarthritis, endotoxemia, toxic shock syndrome, acute and chronic inflammatory diseases, such as the inflammatory reaction induced by endotoxin or inflammatory bowel disease; tuberculosis, atherosclerosis, muscle degeneration, cachexia, psoriatic arthritis, Reiter's syndrome, rheumatoid arthritis, gout, traumatic arthritis, rubella arthritis and acute synovitis. Recent evidence also links IL-1 activity to diabetes.

Interleukin-1 has been demonstrated to mediate a variety of biological activities thought to be important in immunoregulation and other physiological conditions. [See, e.g., Dinarello et al., Rev. Infect. Disease, 6, 51 (1984)]. The known biological activities of IL-1 include the activation of T helper cells, induction of fever, stimulation of prostaglandin or collagenase production, neutrophil chemotaxis, induction of acute phase proteins and the suppression of plasma iron levels.

Excessive or unregulated tumor necrosis factor (TNF) production or activity has been implicated in mediating or exacerbating rheumatoid arthritis, rheumatoid spondylitis, osteoarthritis, gouty arthritis, and other arthritic conditions, sepsis, septic shock, endotoxic shock, gram negative sepsis, toxic shock syndrome,

adult respiratory distress syndrome, cerebral malaria, chronic pulmonary inflammatory disease, silicosis, pulmonary sarcosis, bone resorption diseases, reperfusion injury, graft v. host rejection, allograft rejections, fever and myalgia due to infection, cachexia secondary to infection or malignancy, cachexia secondary to acquired immune deficiency syndrome (AIDS), AIDS related complex (ARC), keloid formation, scar tissue formation, Crohn's disease, ulcerative colitis and pyresis.

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Monokines, such as TNF, have also been shown to activate HIV replication in monocytes and/or macrophages [See Poli, et al., Proc. Natl. Acad. Sci., 87:782-784 (1990)], therefore, inhibition of monokine production or activity aids in limiting HIV progression. TNF has been implicated in various roles with other viral infections, such as the cytomegalovirus (CMV), influenza virus and the herpes virus.

Interleukin-6 (IL-6) is a cytokine effecting the immune system and hematopoiesis. It is produced by several mammalian cell types in response to agents such as IL-1, and is correlated with disease states such as angiofollicular lymphoid hyperplasia.

Interleukin-8 (IL-8) is a chemotactic factor first identified and characterized in 1987. Many different names have been applied to IL-8, such as neutrophil attractant/activation protein-1 (NAP-1), monocyte derived neutrophil chemotactic factor (MDNCF), neutrophil activating factor (NAF), and T-cell lymphocyte chemotactic factor. Like IL-1, IL-8 is produced by several cell types, including mononuclear cells, fibroblasts, endothelial cells and ketainocytes. Its production is induced by IL-1, TNF and by lipopolysaccharide (LPS). IL-8 stimulates a number of cellular functions in vitro. It is a chemoattractant for neutrophils, T-lymphocytes and basophils. It induces histamine release from basophils. It causes lysozomal enzyme release and respiratory burst from neutrophils, and it has been shown to increase the surface expression of Mac-1 (CD11b/CD 18) on neutrophils without de novo protein synthesis.

There remains a need for compounds which are useful in treating cytokine mediated diseases, and as such, inhibit, suppress or antagonize the production or activity of cytokines such as IL-1, IL-6, IL-8 and TNF.

Parasitic protozoa are responsible for a wide variety of infections in man and animals. Many of the diseases are life threatening to the host, and in animal husbandry, can cause considerable economic loss. For example, malaria remains a significant health threat to humans despite massive international attempts to eradicate the disease; trypanosomiasis such as Chagas disease caused by *Trypanosoma cruzi*

and African sleeping sickness caused by *T. brucei* are not uncommon in South America and Africa, respectively; and opportunistic infections in immuno-compromised hosts caused by *Pneumocystis carinii*, *Toxoplasma gondii*, *Cryptosporidium* sp. are becoming increasingly significant in the developed countries.

Coccidiosis, a widespread disease of domesticated animals, is caused by protozoal infection. In the poultry industry, coccidiosis is responsible for high levels of morbidity and mortality in the bird population and may result in extreme economic losses. The infectious agents are protozoa of the genus *Eimeria*. Some of the most significant avian *Eimeria* species include *E. tenella*, *E. acervulina*, *E. necatrix*, *E. brunetti* and *E. maxima*.

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In some protozoal diseases, such as Chagas disease, there is no satisfactory treatment; in others, drug-resistant strains of the protozoa may develop. A biochemical target of antiprotozoal drugs, cGMP dependent protein kinases (PKG), has been identified, the inhibition of which effectively treats protozoal infections such as coccidiosis and Chagas disease.

cGMP dependent protein kinases catalyze the phosphorylation of specific protein substrates. In the absence of cGMP the activity of these enzymes is very low. Thus, the inhibition of such PKG kinases can be lethal to the organism. There is a need for compounds that treat (or prevent by a subtherapeutic prophalactic dosing) coccidiosis, Chagas disease, and toxoplasmosis. Compounds that inhibit the PKG kinase of the infecting protozoa can be such preventive and treating compounds. Importantly, compounds that selectively inhibit the PKG kinase of the infecting protozoa without inhibiting the PKG kinase of mammalian PKG kinase would be lethal to protozoa while being safe for mammals. Accordingly, there is a need for such selective compounds for the treatment of protozoal infections such as coccidiosis, Chagas disease, and toxoplasmosis.

International Patent Publication Nos. WO 99/51233, WO 99/51232, WO 97/21704, WO 97/21703, and WO 00/04013 describe fused heteroaryl compounds that are antagonists of gonadotropin releasing hormone. International Patent Publication No. WO 96/06840 describes diaryl bicyclic heterocycles as inhibitors of cyclooxygenase-2.

International Patent Publication No. WO 98/22457 describes aryl and heteroaryl substituted fused pyrrole antiinflammatory agents. International Patent Publication No. WO 01/22965 describes substituted imidazoles having cytokine inhibitory activity. International Patent Publication No. WO 01/34605 describes

substituted 2-aryl-3-(heteroaryl)-imidazo[1,2-a]primidines. International Patent Publication No. WO 01/30778 describes tiazole and imidazo[4,5-b] pyridine compounds. International Patent Publication No. WO 00/63204 describes substituted azoles.

5 The compounds 3-(2-Methylsulfanylpyrimidin-4-yl)-2-(3-trifluoromethylphenyl)imidazo[1,2-a]-pyrimidine:

and 3-(2-Methylsulfonylpyrimidin-4-yl)-2-(3-trifluoromethylphenyl)imidazo[1,2-a]-pyrimidine:

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were described in International Patent Publication No. WO 01/22965 as intermediates in a process to make a substituted imidazole.

SUMMARY OF THE INVENTION

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The present invention relates to compound I of the formula

wherein FusedHet is

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or a pharmaceutically acceptable salt and/or hydrate thereof, or where applicable, a geometric or optical isomer or racemic mixture thereof.

This invention also relates to a pharmaceutical composition that is comprised of a compound of formula I as defined above in combination with a pharmaceutically acceptable carrier.

Also included in the invention is a method of treating a cytokine mediated disease in a mammal, comprising administering to a mammalian patient in need of such treatment an amount of a compound of formula I which is effective to treat the cytokine mediated disease.

The invention includes a method of treating a protozoal disease in a mammel or bird, comprising administering to a mammalian or avian patient in need of such treatment an amount of a compound of formula I which is effective to treat the protozoal disease. Further, the invention includes a method of preventing a protozoal disease in a mammel or bird, comprising administering to a mammalian or avian patient in need of such treatment a prophalactic amount of a compound of formula I which is effective to prevent the protozoal disease.

DETAILED DESCRIPTION OF THE INVENTION

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The present invention relates to compounds represented by formula (I):

(I)

or a pharmaceutically acceptable salt or hydrate thereof, wherein FusedHet is

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$$R^3$$
, or

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R¹ is H,

 $-C_{1-6}$ alkyl,

 $-C(O)(C_{1-6}alkyl),$

-C(O)-C₁-6alkyl-aryl,

-C₀₋₄alkyl-aryl,

-C₀₋₄alkyl-indanyl,

-C₀₋₄alkyl-imidazolyl,

-C₀₋₄alkyl-thiazolyl,

-C₀-4alkyl-pyrazolyl,

-C₀₋₄alkyl-oxadiazolyl,

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-C₀-4alkyl-C₃-6cycloalkyl,

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-C<sub>0</sub>-4alkyl-C<sub>1</sub>-4alkoxy,
                                   -C_{1-4}alkyl-N(C_{0-4}alkyl)(-C_{0-4}alkyl),
                                   -C<sub>1</sub>-4alkyl-N(-C<sub>0</sub>-4alkyl)-CO-C<sub>1</sub>-4alkoxy,
                                   -C<sub>1-4</sub>alkyl-piperadinyl,
  5
                                   -C<sub>0-4</sub>alkyl-triazolyl,
                                   -C<sub>1-4</sub>alkyl-imidazothiazolyl.
                                  -C<sub>1</sub>-4alkyl-benzimidazolyl,
                                  -C<sub>1</sub>-4alkyl-benzothiazolyl,
                                  -C<sub>1</sub>-4alkyl-benzotetrahydrofuranyl,
10
                                  -C<sub>1</sub>-4alkyl-benzodioxolyl,
                                  -C1_4alkyl-(heterocycloC4O2alkyl),
                                  -C1-4alkyl-(heterocycloC5O1alkyl),
                                  -C<sub>1-4</sub>alkyl-tetrahydrofuran, or
                                  -C<sub>1</sub>-4alkyl-oxetanyl;
15
                           R^{11} is H or -C_{1-6}alkyl;
                           or R1 and R11, together with the N to which they are attached, form
       a morpholinyl;
                           R2, R21, R22 each independently is H, halogen, or -C1-4alkyl;
                           R^3 is H.
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                              -C<sub>1-4</sub>alkyl,
                                  -C3-6cycloalkyl,
                                  -C<sub>1-4</sub>alkyl-aryl,
                                  -C<sub>1-4</sub>alkyl-azetidinyl,
                                  -C1-4alkyl-azetidinyl-CO-C0-4alkyl-N(C0-4alkyl)(C0-4alkyl),
25
                                  -C<sub>1-4</sub>alkyl-pyrrolidinyl,
                                  -C<sub>1-4</sub>alkyl-piperidinyl,
                                  -C1_4alkyl-morpholinyl,
                                  -C_{0-4}alkyl-N(C_{0-4}alkyl)(C_{0-4}alkyl),
                                 -C_{0-4}alkyl-N(C_{0-4}alkyl)(C_{0-4}alkyl-C_{1-4}alkoxy),
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                                 -C0-4alkyl-N(C0-4alkyl-C1-4alkoxy)(C0-4alkyl-C1-4alkoxy),
                                 -C_{1-4}alkyl-N(C_{0-4}alkyl)-(C_{1-4}alkyl)-aryl,
                                 -C1-4alkyl-N(C0-4alkyl)-C1-4alkyl-tetrahydrofuranyl,
                                 -C1-4alkyl-N(C0-4alkyl)-C1-4alkyl-azetidinyl,
                                 -C_{1-4}alkyl-N(C_{0-4}alkyl)-C_{1-4}alkyl-N(C_{0-4}alkyl)(C_{0-4}alkyl),
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		- C_{1-4} alkyl- $N(C_{0-4}$ alkyl)- C_{1-4} alkyl- $N(C_{0-4}$ alkyl)- C_{0-4} alkyl-
	SO ₂ C ₁₋₄ alkyl),	
		-CO-N(C ₀₋₄ alkyl)-C ₁₋₄ alkyl-aryl,
		-CO-N(C ₀₋₄ alkyl)-C ₁₋₄ alkyl-N(C ₀₋₄ alkyl)(C ₀₋₄ alkyl),
5		-C ₀₋₄ alkyl-CO-C ₀₋₄ alkyl,
		-C ₀ -4alkyl-CO-C ₀ -4alkoxy,
		-C ₀₋₄ alkyl-CO-N(C ₀₋₄ alkyl)-C ₁₋₄ alkyl-C ₁₋₄ alkoxy,
		-C ₀₋₄ alkyl-CO-N(C ₀₋₄ alkyl)-C ₁₋₄ alkyl-aryl,
		-C ₀₋₄ alkyl-CO-piperidinyl,
10		-C1-4alkyl-N(C0-4alkyl)-C0-4alkyl-CO-C0-4alkyl-N(C0-
	4alkyl)(C0_4alkyl),	
		-C ₀₋₄ alkyl-CO-C ₀₋₄ alkyl-N(C ₀₋₄ alkyl)(C ₀₋₄ alkyl),
	•	-O-C ₁₋₄ alkyl-aryl,
		-C ₁ -4alkyl-O-C ₁ -4alkyl,
15	•	-C ₀₋₄ alkyl-N(C ₀₋₄ alkyl)-C ₀₋₄ alkyl-CO-C ₀₋₄ alkyl,
		-C ₀₋₄ alkyl-N(C ₀₋₄ alkyl)-C ₀₋₄ alkyl-CO-C ₀₋₄ alkoxy,
		-C ₀₋₄ alkyl-N(C ₀₋₄ alkyl)-C ₀₋₄ alkyl-CO-C ₀₋₄ alkyl-aryl,
		-C ₀₋₄ alkyl-N(C ₀₋₄ alkyl)-C ₀₋₄ alkyl-CO-C ₀₋₄ alkyl(aryl) ₂ ,
		-C0-4alkyl-N(C0-4alkyl)-C0-4alkyl-CO-C0-4alkyl-pyrrolyl,
20	•	-C ₀₋₄ alkyl-N(C ₀₋₄ alkyl)-C ₀₋₄ alkyl-CO-C ₀₋₄ alkyl-
٠	pyrrolidinyl,	
		-C ₀₋₄ alkyl-N(C ₀₋₄ alkyl)-C ₀₋₄ alkyl-CO-C ₀₋₄ alkyl-
	azetidinyl,	
		-C0-4alkyl-N(C0-4alkyl)-C0-4alkyl-CO-C2-4alkenyl-
25	pyrrolidinyl,	
		-C ₀₋₄ alkyl-N(C ₀₋₄ alkyl)-C ₀₋₄ alkyl-CO-C ₀₋₄ alkyl-
	thiophenyl,	
-	•	-C ₀₋₄ alkyl-N(C ₀₋₄ alkyl)-C ₀₋₄ alkyl-CO-C ₂₋₄ alkenyl-
	thiophenyl,	
30		$-C_{0-4}$ alkyl $-N(C_{0-4}$ alkyl) $-C_{0-4}$ alkyl $-C_{0-5}$ $-C_{1-4}$ alkyl $-$ aryl,
•		-C ₀₋₄ alkyl-N(C ₀₋₄ alkyl)-C ₀₋₄ alkyl-CO-C ₃₋₆ cyclolkyl,
		$-C_{0-4}$ alkyl- $N(C_{0-4}$ alkyl) $-C_{0-4}$ alkyl- $CO-O-C_{1-4}$ alkyl-aryl,
		-C ₀₋₄ alkyl-CO-N(C ₀₋₄ alkyl)-C ₀₋₄ alkyl-C ₁₋₄ alkoxy,
		-C ₁₋₄ alkyl-N(C ₀₋₄ alkyl)(-SO ₂ C ₁₋₄ alkyl),

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 $-C_0$ -4alkyl- $N(C_0$ -4alkyl)- C_1 -4alkyl- SO_2C_1 -4alkyl,

-C0-4alkyl-S-C1-4alkyl-aryl,

-C1-4alkyl-PO(C1-4alkoxy)(C1-4alkoxy),

-C₁-4alkyl-azetidinyl-CO-N(C₀-4alkyl)(C₀-4alkyl),

-C₁₋₄alkyl-(heterocycloC₄N₁O₁alkyl),

-C₀₋₄alkyl-CO-(heterocycloC₅N₁alkyl),

-C0-4alkyl-CO-N(C0-4alkyl)-(heterocycloC5N1alkyl),

-C1-4alkyl-(heterocycloC4N2alkyl)-C1-4alkyl,

-C1-4alkyl-(heterocycloC4N2alkyl)-CO-C0-4alkoxy,

-C1-4alkyl-(heterocycloC4N2alkyl)-C1-4alkyl-N(C0-

4alkyl)(C₀₋₄alkyl),

 $-C_1$ -4alkyl-(heterobicycloC5N2alkyl)- C_1 -4alkyl, or

-C1-4alkyl-NH-(heterobicycloC7N1alkyl); and

 R^4 is $-C_{1-6}$ alkyl;

15 wherein any of the above aryl, hetaryl, cycloalkyl, or heterocycloalkyl optionally may be substituted with 1-4 substituents, each substituent independently is halogen, NO2, -CN, -C1-4alkyl, -C0-4alkoxy, -S-C1-4alkyl, or -C0-4alkyl-(CO)-C0-4alkoxy; and any of the above alkyl optionally may be substituted with 1-4 substituents, each substituent independently is halogen, -N3, -CN, 20 -COOH, or -C₀₋₄alkoxy.

This invention also includes a binary compound formed from two compounds of formula (I), as described above, connected together by linking the respective R3 groups of each compound. In one aspect the binary compound is a dimer of two identical compounds of formula (I), as described above.

In one aspect, the compound of this invention is represented by formula (I), or a pharmaceutically acceptable salt or hydrate thereof, wherein FusedHet is

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In a second aspect, the compound of this invention is represented by formula (I), or a pharmaceutically acceptable salt or hydrate thereof, wherein FusedHet is

N N

In a third aspect, the compound of this invention is represented by

formula (I), or a pharmaceutically acceptable salt or hydrate thereof, wherein

FusedHet is

In a fourth aspect, the compound of this invention is represented by formula (I), or a pharmaceutically acceptable salt or hydrate thereof, wherein FusedHet is

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In a fifth aspect, the compound of this invention is represented by formula (I), or a pharmaceutically acceptable salt or hydrate thereof, wherein FusedHet is

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In a sixth aspect, the compound of this invention is represented by formula (I), or a pharmaceutically acceptable salt or hydrate thereof, wherein FusedHet is

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In a seventh aspect, the compound of this invention is represented by formula (I), or a pharmaceutically acceptable salt or hydrate thereof, wherein

15 FusedHet is

In an eighth aspect, the compound of this invention is represented by formula (I), or a pharmaceutically acceptable salt or hydrate thereof, wherein FusedHet is

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In a ninth aspect, the compound of this invention is represented by formula (I), or a pharmaceutically acceptable salt or hydrate thereof, wherein FusedHet is

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In a tenth aspect, the compound of this invention is represented by formula (I), or a pharmaceutically acceptable salt or hydrate thereof, wherein FusedHet is

This invention also relates to a pharmaceutical composition that is comprised of a compound of formula (I) as defined above in combination with a pharmaceutically acceptable carrier.

Also included in the invention is a method of treating a cytokine mediated disease in a mammal, comprising administering to a mammalian patient in need of such treatment an amount of a compound of formula (I), which is effective to treat the cytokine mediated disease.

The invention includes a method of treating a protozoal disease in a mammel, comprising administering to a mammalian patient in need of such treatment an amount of a compound of formula (I), which is effective to treat the protozoal disease. Further, the invention includes a method of preventing a protozoal disease in a mammel, comprising administering to a mammalian patient in need of such treatment a prophalactic amount of a compound of formula (I), which is effective to prevent the protozoal disease.

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Unless otherwise stated or indicated, the following definitions shall apply throughout the specification and claims.

As used herein, "alkyl" as well as other groups having the prefix "alk" such as, for example, alkoxy, alkanoyl, alkenyl, alkynyl and the like, means carbon chains which may be linear or branched or combinations thereof. Examples of alkyl groups include methyl, ethyl, propyl, isopropyl, butyl, sec- and tert-butyl, pentyl, hexyl, heptyl and the like. "Alkenyl", "alkynyl" and other like terms include carbon chains containing at least one unsaturated C-C bond.

The term "cycloalkyl" means carbocycles containing no heteroatoms, and includes mono-, bi- and tricyclic saturated carbocycles, as well as fused ring systems. Such fused ring systems can include one ring that is partially or fully unsaturated such as a benzene ring to form fused ring systems such as benzofused carbocycles. Cycloalkyl includes such fused ring systems as spirofused ring systems. Examples of cycloalkyl include cyclopropyl, cyclobutyl, cyclopentyl, cyclohexyl, decahydronaphthalene, adamantane, indanyl, indenyl, fluorenyl, 1,2,3,4-

tetrahydronaphalene and the like. Similarly, "cycloalkenyl" means carbocycles containing no heteroatoms and at least one non-aromatic C-C double bond, and include mono-, bi- and tricyclic partially saturated carbocycles, as well as benzofused cycloalkenes. Examples of cycloalkenyl include cyclohexenyl, indenyl, and the like.

The term "aryl" means an aromatic substituent which is a single ring or multiple rings fused together. When formed of multiple rings, at least one of the

constituent rings is aromatic. The preferred aryl substituents are phenyl and napthyl groups.

The term "cycloalkyloxy" unless specifically stated otherwise includes a cycloalkyl group connected by a short C₁-C₂alkyl length to the oxy connecting atom.

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The term "C0-C6alkyl" includes alkyls containing 6, 5, 4, 3, 2, 1, or no carbon atoms. A terminal alkyl with no carbon atoms is a hydrogen atom. A bridging alkyl with no carbon atoms is a direct bond. It is understood that, for the purposes of substitution, an alkyl with no carbon atoms has no substituents and takes no substitution. The term "-C0-4alkoxy" is -OH for -C0alkoxy.

The term "hetero" unless specifically stated otherwise includes one or more O, S, or N atoms. For example, heterocycloalkyl ("heterocycle") and heteroaryl include ring systems that contain one or more O, S, or N atoms in the ring, including mixtures of such atoms. The hetero atoms replace ring carbon atoms. Thus, for example, a heterocycloC5alkyl is a five member ring containing from 5 to no carbon atoms. However, the heteroatoms can be specified. Thus, a heterocycloC4N1O1alkyl is a six member saturated ring containing 4 carbon atoms, 1 nitrogen atom, and 1 oxygen atom. Similar notation is used for heterobicycloclkyls.

Generally, unless otherwise stated, "heterocycle" is a 3- to 7-membered non-aromatic ring containing 1-4 heteroatoms selected from N, O and S(O)m, which may be optionally fused to a benzene ring, and in which up to three additional carbon atoms may be replaced by said heteroatoms. When three heteroatoms are present in the heterocycle, they are not all linked together. Examples of heterocycle include oxiranyl, aziridinyl, azetidinyl, oxetanyl, tetrahydrofuranyl, tetrahydropyranyl, tetrahydrothienyl including sulfoxide and sulfones thereof, 2,3- and 2,5-dihydrofuranyl, 1,3-dioxanyl, 1,3-dioxolanyl, pyrrolidinyl, imidazolinyl, imidazolinyl, piperidinyl, morpholinyl, thiomorpholinyl, piperazinyl, benzoxazinyl, 2,3-dihydrobenzofuranyl 1,2,3,4-tetrahydroisoquinolinyl, 1,2,3,4-tetrahydroquinolinyl.

"Heteroaryl" is a mono-or bicyclic aromatic ring containing from 1 to 6 heteroatoms independently selected from N, O and S wherein each ring has five or six ring atoms. Examples of heteroaryl include pyridyl, pyrimidinyl, pyrrolyl, furyl, thienyl, imidazolyl, thiazolyl, thiadiazolyl, triazolyl, tetrazolyl, oxadiazolyl, oxazolyl, imidazolidinyl, pyrazolyl, isoxazolyl, benzothiadiazolyl, indolyl, indolinyl, benzodioxolyl, benzodioxanyl, benzothiophenyl, benzofuranyl, benzimidazolyl,

benzisoxazolyl, benzothiazolyl, quinolinyl, benzotriazolyl, benzoxazolyl, purinyl, furopyridine and thienopyridine.

The term "amine" unless specifically stated otherwise includes primary, secondary and tertiary amines.

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The term "halogen" or "halo" is intended to include fluorine, chlorine, bromine and iodine.

The term "optionally substituted" is intended to include both substituted and unsubstituted. Thus, for example, optionally substituted aryl could represent a pentafluorophenyl or a phenyl ring. Further, optionally substituted multiple moieties such as, for example, alkylaryl are intended to mean that the aryl and the aryl groups are optionally substituted. If only one of the multiple moieties is optionally substituted then it will be specifically recited such as "an alkylaryl, the aryl optionally substituted with halogen or hydroxyl."

The term "composition", as in pharmaceutical composition, is intended to encompass a product comprising the active ingredient(s), and the inert ingredient(s) that make up the carrier, as well as any product which results, directly or indirectly, from combination, complexation or aggregation of any two or more of the ingredients, or from dissociation of one or more of the ingredients, or from other types of reactions or interactions of one or more of the ingredients. Accordingly, the pharmaceutical compositions of the present invention encompass any composition made by admixing a compound of the present invention and a pharmaceutically acceptable carrier.

Compounds described herein contain one or more double bonds and may thus give rise to cis/trans isomers as well as other conformational isomers. The present invention includes all such possible isomers as well as mixtures of such isomers.

Compounds described herein can contain one or more asymmetric centers and may thus give rise to diastereomers and optical isomers. The present invention includes all such possible diastereomers as well as their racemic mixtures, their substantially pure resolved enantiomers, all possible geometric isomers, and pharmaceutically acceptable salts thereof. The above Formula I is shown without a definitive stereochemistry at certain positions. The present invention includes all stereoisomers of Formula I and pharmaceutically acceptable salts thereof. Further, mixtures of stereoisomers as well as isolated specific stereoisomers are also included. During the course of the synthetic procedures used to prepare such compounds, or in

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using racemization or epimerization procedures known to those skilled in the art, the products of such procedures can be a mixture of stereoisomers.

The term "pharmaceutically acceptable salts" refers to salts prepared from pharmaceutically acceptable non-toxic bases or acids. When the compound of the present invention is acidic, its corresponding salt can be conveniently prepared from pharmaceutically acceptable non-toxic bases, including inorganic bases and organic bases. Salts derived from such inorganic bases include aluminum, ammonium, calcium, copper (ic and ous), ferric, ferrous, lithium, magnesium, manganese (ic and ous), potassium, sodium, zinc and the like salts. Particularly preferred are the ammonium, calcium, magnesium, potassium and sodium salts. Salts 10 derived from pharmaceutically acceptable organic non-toxic bases include salts of primary, secondary, and tertiary amines, as well as cyclic amines and substituted amines such as naturally occurring and synthesized substituted amines. Other pharmaceutically acceptable organic non-toxic bases from which salts can be formed include ion exchange resins such as, for example, arginine, betaine, caffeine, choline, N,N'-dibenzylethylenediamine, diethylamine, 2-diethylaminoethanol, 2dimethylaminoethanol, ethanolamine, ethylenediamine, N-ethylmorpholine, Nethylpiperidine, glucamine, glucosamine, histidine, hydrabamine, isopropylamine, lysine, methylglucamine, morpholine, piperazine, piperidine, polyamine resins, procaine, purines, theobromine, triethylamine, trimethylamine, tripropylamine, tromethamine and the like.

When the compound of the present invention is basic, its corresponding salt can be conveniently prepared from pharmaceutically acceptable non-toxic acids, including inorganic and organic acids. Such acids include, for example, acetic, benzenesulfonic, benzoic, camphorsulfonic, citric, ethanesulfonic, fumaric, gluconic, glutamic, hydrobromic, hydrochloric, isethionic, lactic, maleic, malic, mandelic, methanesulfonic, mucic, nitric, pamoic, pantothenic, phosphoric, succinic, sulfuric, tartaric, p-toluenesulfonic acid and the like. Particularly preferred are citric, hydrobromic, hydrochloric, maleic, phosphoric, sulfuric, and tartaric acids.

The compounds of the present invention may have chiral centers other than those centers whose stereochemistry is depicted in formula I, and therefore may occur as racemates, racemic mixtures and as individual enantiomers or diastereomers, with all such isomeric forms being included in the present invention as well as mixtures thereof. Furthermore, some of the crystalline forms for compounds of the present invention may exist as polymorphs and as such are

intended to be included in the present invention. In addition, some of the compounds of the instant invention may form solvates with water or common organic solvents. Such solvates are encompassed within the scope of this invention.

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The term "TNF mediated disease or disease state" refers to disease states in which TNF plays a role, either by production or increased activity levels of TNF itself, or by causing another cytokine to be released, such as but not limited to IL-1 or IL-6. A disease state in which IL-1, for instance is a major component, and whose production or action, is exacerbated or secreted in response to TNF, would therefore be considered a disease state mediated by TNF.

The term "cytokine" as used herein means any secreted polypeptide that affects the functions of cells and is a molecule which modulates interactions between cells in the immune, inflammatory or hematopoietic response. A cytokine includes, but is not limited to, monokines and lymphokines regardless of which cells produce them. Examples of cytokines include, but are not limited to, Interleukin-1 (IL-1), Interleukin-6 (IL-6), Interleukin-8 (IL-8), Tumor Necrosis Factor-alpha (TNF-α) and Tumor Necrosis Factor-beta (TNF-β).

By the term "cytokine interfering or cytokine suppresive amount" is meant an effective amount of a compound of formula I which will cause a decrease in the *in vivo* activity or level of the cytokine to normal or sub-normal levels, when given to the patient for the prophylaxis or therapeutic treatment of a disease state which is exacerbated by, or caused by, excessive or unregulated cytokine production or activity.

The compounds of formula 1 can be used in the prophylactic or therapeutic treatment of disease states in mammals which are exacerbated or caused by excessive or unregulated cytokines, e.g., IL-1, IL-6, IL-8 or TNF.

Because the compounds of formula I inhibit cytokines, the compounds are useful for treating diseases in which cytokine presence or activity is implicated, such as rheumatoid arthritis, rheumatoid spondylitis, osteoarthritis, gouty arthritis and other arthritic conditions.

The compounds of formula I are useful to treat disease states mediated by excessive or unregulated TNF production or activity. Such diseases include, but are not limited to sepsis, septic shock, endotoxic shock, gram negative sepsis, toxic shock syndrome, adult respiratory distress syndrome, cerebral malaria, chronic pulmonary inflammatory disease, silicosis, pulmonary sarcoidosis, bone resorption

diseases, such as osteoporosis, reperfusion injury, graft v. host rejection, allograft rejection, fever, myalgia due to infection, cachexia secondary to infection or malignancy, cachexia secondary to acquired immune deficiency syndrome (AIDS), AIDS, ARC (AIDs related complex), keloid formation, scar tissue formation, Crohn's disease, ulcerative colitis, pyresis, AIDS and other viral infections, such as cytomegalovirus (CMV), influenza virus, and the herpes family of viruses such as Herpes Zoster or Simplex I and II.

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The compounds of formula I are also useful topically in the treatment of inflammation such as in the treatment of rheumatoid arthritis, rheumatoid spondylitis, osteoarthritis, gouty arthritis and other arthritic conditions; inflamed joints, eczema, psoriasis or other inflammatory skin conditions such as sunburn; inflammatory eye conditions including conjunctivitis; pyresis, pain and other conditions associated with inflammation.

The compounds of formula I are also useful in treating diseases characterized by excessive IL-8 activity. These disease states include psoriasis, inflammatory bowel disease, asthma, cardiac and renal reperfusion injury, adult respiratory distress syndrome, thrombosis and glomerulonephritis.

The invention thus includes a method of treating psoriasis, inflammatory bowel disease, asthma, cardiac and renal reperfusion injury, adult respiratory distress syndrome, thrombosis and glomerulonephritis, in a mammal in need of such treatment, which comprises administering to said mammal a compound of formula I in an amount which is effective for treating said disease or condition.

When administered to a patient for the treatment of a disease in which a cytokine or cytokines are implicated, the dosage used can be varied within wide limits, depending upon the type of disease, the age and general condition of the patient, the particular compound administered, the presence or level of toxicity or adverse effects experienced with the drug and other factors. A representative example of a suitable dosage range is from as low as about 0.01mg/kg to as high as about 100mg/kg. However, the dosage administered is generally left to the discretion of the physician.

The methods of treatment can be carried out by delivering the compound of formula I parenterally. The term 'parenteral' as used herein includes intravenous, intramuscular, or intraperitoneal administration. The subcutaneous and intramuscular forms of parenteral administration are generally preferred. The instant

invention can also be carried out by delivering the compound of formula I through subcutaneous, intranasal, intrarectal, transdermal or intravaginal routes.

The compounds of formula I may also be administered by inhalation. By inhalation'is meant intranasal and oral inhalation administration. Appropriate dosage forms for such administration, such as an aerosol formulation or a metered dose inhaler, may be prepared by convention techniques.

The invention also relates to a pharmaceutical composition comprising a compound of formula I and a pharmaceutically acceptable carrier. The compounds of formula I may also be included in pharmaceutical compositions in combination with a second therapeutically active compound.

The pharmaceutical carrier employed may be, for example, either a solid, liquid or gas. Examples of solid carriers include lactose, terra alba, sucrose, talc, gelatin, agar, pectin, acacia, magnesium stearate, stearic acid and the like. Examples of liquid carriers are syrup, peanut oil, olive oil, water and the like. Examples of gaseous carriers include carbon dioxide and nitrogen.

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Similarly, the carrier or diluent may include time delay material well known in the art, such as glyceryl monostearate or glyceryl distearate, alone or with a wax.

A wide variety of pharmaceutical dosage forms can be employed. If a solid dosage is used for oral administration, the preparation can be in the form of a tablet, hard gelatin capsule, troche or lozenge. The amount of solid carrier will vary widely, but generally the amount of the present compound will be from about 0.025mg to about 1g with the amount of solid carrier making up the difference to the desired tablet, hard gelatin capsule, troche or lozenge size. Thus, the tablet, hard gelatin capsule, troche or lozenge conveniently would have, for example, 0.025mg, 0.05mg, 0.1mg, 0.5mg, 1mg, 5mg, 10mg, 25mg, 100mg, 250mg, 500mg, or 1000mg of the present compound. The tablet, hard gelatin capsule, troche or lozenge is given conveniently once, twice or three times daily.

When a liquid dosage form is desired for oral administration, the preparation is typically in the form of a syrup, emulsion, soft gelatin capsule, suspension or solution. When a parenteral dosage form is to be employed, the drug may be in solid or liquid form, and may be formulated for administration directly or may be suitable for reconstitution.

Topical dosage forms are also included. Examples of topical dosage forms are solids, liquids and semi-solids. Solids would include dusting powders,

poultices and the like. Liquids include solutions, suspensions and emulsions. Semisolids include creams, ointments, gels and the like.

The amount of a compound of formula I used topically will, of course, vary with the compound chosen, the nature and severity of the condition, and can be varied in accordance with the discretion of the physician. A representative, topical, dose of a compound of formula I is from as low as about 0.01mg to as high as about 2.0g, administered one to four, preferably one to two times daily.

The active ingredient may comprise, for topical administration, conveniently from about 0.001% to about 10% w/w.

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Drops according to the present invention may comprise sterile or non-sterile aqueous or oil solutions or suspensions, and may be prepared by dissolving the active ingredient in a suitable aqueous solution, optionally including a bactericidal and/or fungicidal agent and/or any other suitable preservative, and optionally including a surface active agent. The resulting solution may then be clarified by filtration, transferred to a suitable container which is then sealed and sterilized by autoclaving or maintaining at 98-100°C for half an hour. Alternatively, the solution may be sterilized by filtration and transferred to the container aseptically. Examples of bactericidal and fungicidal agents suitable for inclusion in the drops are phenylmercuric nitrate or acetate (0.002%), benzalkonium chloride (0.01%) and chlorhexidine acetate (0.01%). Suitable solvents for the preparation of an oily solution include glycerol, diluted alcohol and propylene glycol.

Lotions according to the present invention include those suitable for application to the skin or eye. An eye lotion may comprise a sterile aqueous solution optionally containing a bactericide and may be prepared by methods similar to those for the preparation of drops. Lotions or liniments for application to the skin may also include an agent to hasten drying and to cool the skin, such as an alcohol or acetone, and/or a moisturizer such as glycerol or an oil such as castor oil or arachis oil.

Creams, ointments or pastes according to the present invention are semi-solid formulations of the active ingredient for external application. They may be made by mixing the active ingredient in finely-divided or powdered form, alone or in solution or suspension in an aqueous or non-aqueous liquid, with a greasy or non-greasy base. The base may comprise hydrocarbons such as hard, soft or liquid paraffin, glycerol, beeswax, a metallic soap; a mucilage; an oil of natural origin such as almond, corn, arachis, castor or olive oil; wool fat or its derivatives, or a fatty acid such as stearic or oleic acid together with an alcohol such as propylene glycol or

macrogels. The formulation may incorporate any suitable surface active agent such as an anionic, cationic or non-ionic surfactant such as sorbitan esters or polyoxyethylene derivatives thereof. Suspending agents such as natural gums, cellulose derivatives or inorganic materials such as silicas, and other ingredients such as lanolin may also be included.

The ability of compounds of the present invention to inhibit the synthesis or the activity of cytokines can be demonstrated using the following *in vitro* assays.

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BIOLOGICAL ASSAYS

Lipopolysaccharide mediated production of cytokines

Human peripheral blood mononuclear cells (PBMC) are isolated from fresh human blood according to the procedure of Chin and Kostura, J. Immunol., 15 151:5574-5585(1993). Whole blood is collected by sterile venipuncture into 60mL syringes coated with 1.0mL of sodium-heparin (Upjohn, 1000µ/mL) and diluted 1:1 in Hanks Balanced Salt Solution (Gibco). The erythrocytes are separated from the PBMC's by centrifugation on a Ficoll-Hypaque lymphocyte separation media. The 20 PBMC's are washed three times in Hanks Balanced Salt Solution and then resuspended to a final concentration of 2 x 10⁶ cell/mL in RPMI containing 10% fresh autologous human serum, penicillin streptomycin (10µ/mL) and 0.05% DMSO. Lipopolysaccharide (Salmonella type Re545; Sigma Chemicals) is added to the cells to a final concentration of 100ng/mL. An aliquot (0.1mL) of the cells is quickly dispensed into each well of a 96 well plate containing 0.1mL of the test compound, at 25 the appropriate dilution, and are incubated for 24 hours at 37°C in 5% CO₂. At the end of the culture period, cell culture supernatants are assayed for IL-1β, TNF-α, IL-6 and PGE2 production using specific ELISA.

30 IL-1 mediated cytokine production

Human peripheral blood mononuclear cells are isolated from fresh human blood according to the procedure of Chin and Kostura, *J. Immunol.*, 151:5574-5585(1993). Whole blood is collected by sterile venipuncture into 60mL syringes coated with 1.0mL of sodium-heparin (Upjohn, 1000µ/mL) and diluted 1:1 in Hanks

Balanced Salt Solution (Gibco). The erythrocytes are separated from the PBMC's by centrifugation on a Ficoll-Hypaque lymphocyte separation media. The PBMC's are washed three times in Hanks Balanced Salt Solution and then resuspended to a final concentration of 2 x 10⁶ cell/mL in RPMI containing 10% fresh autologous human serum, penicillin streptomycin (10μ/mL) and 0.05% DMSO. Endotoxin free recombinant human IL-1b is then added to a final concentration of 50 pMolar. An aliquot (0.1mL) of the cells is quickly dispensed into each well of a 96 well plate containing 0.1mL of the compound at the appropriate dilution and incubated for 24 hours at 37°C in 5% CO₂. At the end of the culture period, cell culture supernatants are assayed for TNF-a, IL-6 and PGE2 synthesis using specific ELISA.

Determination of IL-1β, TNF-α, IL-6 and Prostanoid Production from LPS or IL-1 Stimulated PBMC's

15 IL-1β ELISA

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Human IL-1 β can be detected in cell-culture supernatants or whole blood with the following specific trapping ELISA. 96 well plastic plates (Immulon 4; Dynatech) are coated for 12 hours at 4°C with 1mg/mL protein-A affinity chromatography purified mouse anti-human IL-1β monoclonal antibody (purchased as an ascites preparation from LAO Enterprise, Gaithersburg Maryland.) diluted in 20 Dulbecco's phosphate-buffered saline (-MgCl₂, -CaCl₂). The plates are washed with PBS-Tween (Kirkegaard and Perry) then blocked with 1% BSA diluent and blocking solution (Kirkegaard and Perry) for 60 minutes at room temperature followed by washing with PBS Tween. IL-1β standards are prepared from purified recombinant 25 IL-1β produced from E. coli. The highest concentration begins at 10ng/mL followed by 11 two-fold serial dilutions. For detection of IL-1β from cell culture supernatants or blood plasma, 10 - 25mL of supernatant is added to each test well with 75-90mL of PBS Tween. Samples are incubated at room temperature for 2 hours then washed 6 times with PBS Tween on an automated plate washer (Dennly). Rabbit anti-human 30 IL-1β polyclonal antisera diluted 1:500 in PBS-Tween is added to the plate and incubated for 1 hour at room temperature followed by six washes with PBS-Tween, Detection of bound rabbit anti-IL-1β IgG is accomplished with Fab' fragments of Goat anti-rabbit IgG-horseradish peroxidase conjugate (Accurate Scientific) diluted 1:10,000 in PBS-Tween. Peroxidase activity was determined using TMB peroxidase

substrate kit (Kirkegaard and Perry) with quantitation of color intensity on a 96-well plate Molecular Devices spectrophotometer set to determine absorbance at 450 nM. Samples are evaluated using a standard curve of absorbance versus concentration. Four-parameter logistics analysis generally is used to fit data and obtain concentrations of unknown compounds.

TNF-α ELISA

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Immulon 4 (Dynatech) 96-well plastic plates are coated with a 0.5mg/mL solution of mouse anti-human TNF-a monoclonal antibody. The secondary antibody is a 1:2500 dilution of a rabbit anti-human TNF-α polyclonal serum purchased from Genzyme. All other operations are identical to those described above for IL-1β. The standards are prepared in PBS-Tween + 10% FBS or HS. Eleven two-fold dilutions are made beginning at 20ng/mL TNF-α.

15 IL-6 ELISA

Levels of secreted human IL-6 are also determined by specific trapping ELISA as described previously in Chin and Kostura, *J. Immunol.*, 151:5574-5585(1993). (Dynatech) ELISA plates are coated with mouse anti-human IL-6 monoclonal antibody diluted to 0.5mg/mL in PBS. The secondary antibody, a rabbit anti-human IL-6 polyclonal antiserum, is diluted 1:5000 with PBS-Tween. All other operations are identical to those described above for IL-1β. The standards are prepared in PBS-Tween + 10% FBS or HS. Eleven two-fold dilutions are made beginning at 50ng/mL IL-6.

25 PGE₂ production

Prostaglandin E2 is detected in cell culture supernatants from LPS or IL-1 stimulated PBMC's using a commercially available enzyme immunoassay. The assay purchased from the Cayman Chemical (Catalogue No. 514010) and is run exactly according to the manufacturers instructions.

Interleukin-8 (IL-8)

The present compounds can also be assayed for IL-8 inhibitory activity as discussed below. Primary human umbilical cord endothelial cells (HUVEC) (Cell Systems, Kirkland, Wa) are maintained in culture medium supplemented with 15% fetal bovine serum and 1% CS-HBGF consisting of aFGF

and heparin. The cells are then diluted 20-fold before being plated (250 μ L) into gelatin coated 96-well plates. Prior to use, culture medium is replaced with fresh medium (200 μ L). Buffer or test compound (25 μ L, at appropriate concentrations) is then added to each well in quadruplicate wells and the plates incubated for 6h in a humidified incubator at 37°C in an atmosphere of 5% CO2. At the end of the incubation period, supernatant is removed and assayed for IL-8 concentration using an IL-8 ELISA kit obtained from R&D Systems (Minneapolis, MN). All data is presented as mean value (ng/mL) of multiple samples based on the standard curve. IC50 values where appropriate are generated by non-linear regression analysis.

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The compounds of this invention, in the above functional activity assay, suppress TNF-α in monocytes with IC50 of less than 5μM. Advantageously, the IC50 should be less than 3µM. Even more advantaeously, the IC50 should be less than 1µM. Still more advantageously, the IC50 should be less than 0.1µM.

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Further, in the other assays, the results from the present compounds are better than 5μM. Advantageously, the IC50 results should be less than 3μM. Even more advantageously, the IC50 should be less than 1µM. Still more advantageously, the IC50 should be less than 0.1 µM.

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The ability of compounds of the present invention to inhibit the activity of protozoa can be demonstrated using the following assays.

Anticoccidiosis Assay.

25 One-day-old White Leghorn chickens are obtained from a commercial hatchery and acclimated in a holding room. At three days of age the test animals are selected by weight, wingbanded, and randomly placed on medicated or control diets for the duration of the experiment. One or two replicates of two birds are utilized per treatment. Following 24h premedication, in each replicate one bird is infected with Eimeria acervulina, the other bird is infected with E. tenella. Both strains of Eimeria 30 are sensitive to all anticoccidial products, and have been maintained in laboratory conditions for over 25 years. The inocula consist of sporulated oocysts in tap water suspensions, administered at a dose rate of 0.25mL per bird. The inocula levels are selected by previous dose titrations to provide a low to moderate level of infection.

35 The E. acervulina portion of the experiment is terminated on Day 5, the E. tenella on

Day 6 post infection. The measured parameters are weight gain, feed consumption and oocyst production. *E. tenella* lesion scores are also recorded for background information. Treatments which provide at least 80% reduction in oocyst production are considered active, those with 50-79% are considered partially active, and those with <50% are considered inactive. The same numerical categories in weight gain and feed consumption differentiate among treatments with good, fair or poor productivity.

PKG Catalytic Assay

Kinase activity was detected using a peptide substrate and [³³P]-ATP. An aliquot containing enzyme (1μl) was mixed with a reaction mix (10μl) whose composition is as follows: 25mM HEPES pH 7.4, 10mM MgCl₂, 20mM β-glycerophosphate, 5mM β mercaptoethanol, 10μM cGMP, 1mg/mL BSA, 400μM kemptide, 2μM [³³P]ATP (0.1mCi/ml). The reaction was allowed to proceed for 1 hour at room temperature prior to addition of phosphoric acid to a final concentration of 2.5mM. Labeled peptide was captured on filters using either P81 filters or on Millipore 96-well plates, MAPH-NOB (Millipore). In both cases filters were washed with 75mM phosphoric acid, dried and [³³P]-ATP detected using scintillation counting.

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Enzyme assay and data analysis

The peptide substrate biotinyl-ε-aminocaproyl-GRTGRRNSI-OH was synthesized in house by standard methods. PET-cGMP, 1-NH₂-cGMP, 8-APT-cGMP, and 8-NBD-cGMP were obtained from Biolog Life Science Institute (Bremen, FRG), while 8-Br-cGMP came from Biomol Research Laboratories and 8-pCPT-cGMP came from Calbiochem. Bovine PKG was obtained commercially; recombinant isoform Iα (Genbank Accession No. X16086) was purchased from Calbiochem, while native Iα enzyme was purchased from Promega.

The kinase assay was performed in a 50μ L reaction volume containing 25mM HEPES (pH 7.0), 10mM MgCl₂, 20mM beta-glycerophosphate, 1mM DTT, 0.1mg/mL bovine serum albumin, 20μ M ATP, 20μ M peptide substrate and 2.5μ Ci [gamma-³³P]ATP (Amersham). Cyclic nucleotide was serially diluted in buffer before adding 5μ L of each concentration into 40μ L of the assay mix. The reaction was initiated with 5μ L of enzyme (or buffer for the background) and incubated for 30 minutes in a heating block at 30°C. The assays were terminated by the addition of

 25μ L 8M guanidine-HCl solution (Pierce) before spotting 15μ L onto a SAM² streptavidin membrane (Promega). The membrane was washed twice with 1M NaCl and twice with 1M NaCl + 1% H₃PO₄ on a rotating mixer for 20 minutes. The membrane was then rinsed successively with water and ethanol and dried under a heat lamp.

The individual assays were then separated, placed in scintillation vials containing 2mL of Ultima Gold cocktail (Packard), and counted in a Packard TriCarb 2500 liquid scintillation counter. The amount of enzyme was adjusted to give between 10,000 and 140,000 cpm when maximally activated; substrate turnover was less than 10% in all cases. The concentration of Et-PKG varied between 0.26 and 3.4 µg/mL for cGMP titrations and between 7 and 25µg/mL for 8-NBD-cGMP titrations, depending on the activity of the enzyme form used. Assays with bovine PKG used 0.059 µg/mL recombinant or 0.034 µg/mL native enzyme with both activators. After subtracting the appropriate background for each assay point, titrations were fit to the following modified Hill equation using Kaleidagraph (Synergy Software):

$$V_A = V_0 + (V_{max} - V_0)/(1 + (K_A/[A])^h)$$

 V_A is the observed velocity at concentration [A] of cyclic nucleotide, V_0 is the velocity in the absence of activator, V_{max} is the velocity of the maximally activated enzyme, K_A is the concentration for half maximal activation, and h is the Hill coefficient. The activation parameters are determined from a curve fit.

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cGMP-Agarose Affinity Chromatography-

Purification of PKG enzyme was performed as follows. Chromatography on cGMP-agarose was performed according to the manufacturers instructions (Biolog, A019). Briefly, a 0.6mL column was equilibrated with Buffer G (50mM HEPES pH 7.4, 10% glycerol, 10mM sodium fluoride, 0.1mM sodium orthovanadate, 1mM EDTA). The sample (crude S100 extract or purified protein) was mixed with an equal volume of Buffer G and applied to the column; the column was then washed with 10mL of the same buffer. The column was then washed with

10mL of Buffer G containing 1mM GMP. PKG was then eluted with 10mL of Buffer G containing 15mM cGMP.

In the above assays, the compounds show selectivity, with inhibition of the parasitic enzyme with negligible inhibition of the host enzyme. Thus, it is advantageous that the parasite PKG enzyme IC50 be less than 0.5μM while the host PKG enzyme IC50 be greater than 1μM. It is more advantageous that the host PKG IC50 be greater than 5μM. It is also more advantageous that the parasite PKG enzyme IC50 be less than 0.1μM. It is even more advantageous that the parasite PKG enzyme IC50 be less than 50nM, and particularly advantageous that the parasite PKG enzyme IC50 be less than 10nM.

15 Utility

The (pyrimidyl)(phenyl)substituted fused heteroaryl compounds of the present invention are useful as antiprotozoal agents. As such, they may be used in the treatment and prevention of protozoal diseases in human and animals, including poultry. Examples of protozoal diseases against which compounds of formula I may 20 be used, and their respective causative pathogens, include: 1) amoebiasis (Dientamoeba sp., Entamoeba histolytica); 2) giardiasis (Giardia lamblia); 3) malaria (Plasmodium species including P. vivax, P. falciparum, P. malariae and P. ovale); 4) leishmaniasis (Leishmania species including L. donovani, L. tropica, L. mexicana, and L. braziliensis); 5) trypanosomiasis and Chagas disease (Trypanosoma species 25 including T. brucei, T. theileri, T. rhodesiense, T. gambiense, T. evansi, T. equiperdum, T. equinum, T. congolense, T. vivax and T. cruzi); 6) toxoplasmosis (Toxoplasma gondii); 7) babesiosis (Babesia sp.); 8) cryptosporidiosis (Cryptosporidium sp.); 9) dysentery (Balantidium coli); 10) vaginitis (Trichomonas species including T.vaginitis, and Tritrichomonas foetus); 11) coccidiosis (Eimeria species including E. tenella, E. necatrix, E. acervulina, E. maxima and E. brunetti, E. 30 mitis, E. bovis, E. melagramatis, and Isospora sp.); 12) enterohepatitis (Histomonas gallinarum), and 13) infections caused by Anaplasma sp., Besnoitia sp., Leucocytozoan sp., Microsporidia sp., Sarcocystis sp., Theileria sp., and Pneumocystis carinii.

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Dose Range:

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Compounds of formula I may be administered to a host in need of treatment in a manner similar to that used for other antiprotozoal agents; for example, they may be administered parenterally, orally, topically, or rectally. The dosage to be administered will vary according to the particular compound used, the infectious organism involved, the particular host, the severity of the disease, physical condition of the host, and the selected route of administration; the appropriate dosage can be readily determined by a person skilled in the art.

For the treatment of protozoal diseases in humans, the oral dosage may range from 1mg/kg to 1000mg/kg; and the parenteral dosage may range from 0.5mg/kg to 500mg/kg. For veterinary therapeutic use, the oral dosage may range from 1mg/kg to 1000mg/kg; and the parenteral dosage may range from 0.5mg/kg to 500mg/kg. For prophylactic use in humans, the oral dosage may range from 1mg/kg to 1000mg/kg; and the parenteral dosage may range from 0.5mg/kg to 500mg/kg.

Thus, the tablet, hard gelatin capsule, troche or lozenge conveniently would have, for example, 0.1mg, 0.5mg, 1mg, 5mg, 10mg, 25mg, 100mg, 250mg, 500mg, or 1000mg of the present compound. The tablet, hard gelatin capsule, troche or lozenge is given conveniently once, twice or three times daily.

For prophylactic use in animal, the oral dosage may range from 1mg/kg to 1000mg/kg; and the parenteral dosage may range from 0.5mg/kg to 500mg/kg. For use as an anticoccidial agent, particularly in poultry, the compound may be administered in the animals' feed or drinking water in accordance with common practice in the poultry industry and as described below.

The compositions of the present invention comprises a compound of formula I and an inert carrier. The compositions may be in the form of pharmaceutical compositions for human and veterinary usage, or in the form of feed composition for the control of coccidiosis in poultry.

The pharmaceutical compositions of the present invention comprise a compound of formula I as an active ingredient, and may also contain a physiologically acceptable carrier and optionally other therapeutic ingredients. The compositions include compositions suitable for oral, rectal, topical, and parenteral (including subcutaneous, intramuscular, and intravenous) administrations, although the most suitable route in any given case will depend on the particular host, and nature and severity of the conditions for which the active ingredient is being administered. The

pharmaceutical compositions may be conveniently presented in unit dosage form and prepared by any of the methods well-known in the art of pharmacy.

In practical use, compounds of formula I can be combined as the active ingredient in intimate admixture with a pharmaceutical carrier according to conventional pharmaceutical compounding techniques. The carrier may take a wide variety of forms depending on the form of preparation desired for administration, e.g., oral or parenteral (including intravenous).

In preparing the compositions for oral dosage form, any of the usual pharmaceutical media may be employed. For example, in the case of oral liquid preparations such as suspensions, elixirs and solutions, water, glycols, oils, alcohols, flavoring agents, preservatives, coloring agents and the like may be used; or in the case of oral solid preparations such as powders, capsules and tablets, carriers such as starches, sugars, microcrystalline cellulose, diluents, granulating agents, lubricants, binders, disintegrating agents, and the like may be included. Because of their ease of administration, tablets and capsules represent the most advantageous oral dosage unit form in which case solid pharmaceutical carriers are obviously employed. If desired, tablets may be coated by standard aqueous or nonaqueous techniques. In addition to the common dosage forms set out above, compounds of formula I may also be administered by controlled release means and/or delivery devices.

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Pharmaceutical compositions of the present invention suitable for oral administration may be presented as discrete units such as capsules, cachets or tablets each containing a predetermined amount of the active ingredient, as a powder or granules or as a solution or a suspension in an aqueous liquid, a non-aqueous liquid, an oil-in-water emulsion or a water-in-oil liquid emulsion. Such compositions may be prepared by any of the methods of pharmacy but all methods include the step of bringing into association the active ingredient with the carrier which constitutes one or more necessary ingredients. In general, the compositions are prepared by uniformly and intimately admixing the active ingredient with liquid carriers or finely divided solid carriers or both, and then, if necessary, shaping the product into the desired presentation. For example, a tablet may be prepared by compression or molding, optionally with one or more accessory ingredients. Compressed tablets may be prepared by compressing, in a suitable machine, the active ingredient in a freeflowing form such as powder or granules, optionally mixed with a binder, lubricant, inert diluent, surface active or dispersing agent. Molded tablets may be made by molding in a suitable machine, a mixture of the powdered compound moistened with

an inert liquid diluent. Desirably, each tablet contains from about 1mg to about 500mg of the active ingredient and each cachet or capsule contains from about 1 to about 500mg of the active ingredient.

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Pharmaceutical compositions of the present invention suitable for parenteral administration may be prepared as solutions or suspensions of these active compounds in water suitably mixed with a surfactant such as hydroxypropylcellulose. Dispersions can also be prepared in glycerol, liquid polyethylene glycols, and mixtures thereof in oils. Under ordinary conditions of storage and use, these preparations contain a preservative to prevent the growth of microorganisms.

The pharmaceutical forms suitable for injectable use include sterile aqueous solutions or dispersions and sterile powders for the extemporaneous preparation of sterile injectable solutions or dispersions. In all cases, the form must be sterile and must be fluid to the extent that easy syringability exists. It must be stable under the conditions of manufacture and storage and must be preserved against the contaminating action of microorganisms such as bacteria and fungi. The carrier can be a solvent or dispersion medium containing, for example, water, ethanol, polyol (e.g. glycerol, propylene glycol and liquid polyethylene glycol), suitable mixtures thereof, and vegetable oils.

Suitable topical formulations include transdermal devices, aerosols, creams, ointments, lotions, dusting powders, and the like. These formulations may be prepared via conventional methods containing the active ingredient. To illustrate, a cream or ointment is prepared by mixing sufficient quantities of hydrophilic material and water, containing from about 5-10% by weight of the compound, in sufficient quantities to produce a cream or ointment having the desired consistency.

Pharmaceutical compositions suitable for rectal administration wherein the carrier is a solid may be presented as unit dose suppositories. Suitable carriers include cocoa butter and other materials commonly used in the art, and the suppositories may be conveniently formed by admixture of the combination with the softened or melted carrier(s) followed by chilling and shaping molds.

It should be understood that in addition to the aforementioned carrier ingredients the pharmaceutical formulations described above may include, as appropriate, one or more additional carrier ingredients such as diluents, buffers, flavoring agents, binders, surface-active agents, thickeners, lubricants, preservatives (including anti-oxidants) and the like, and substances included for the purpose of rendering the formulation isotonic with the blood of the intended recipient.

For use in the management of coccidiosis in poultry, a compound of formula I may be conveniently administered as a component of a feed composition. Suitable poultry feed composition will typically contain from about 1 ppm to about 1000 ppm, or from about 0.0005% to about 0.05% percent, by weight of a compound of formula I. The optimum levels will naturally vary with the species of *Eimeria* involved, and can be readily determined by one skilled in the art.

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In the preparation of poultry feed, a compound of formula I may be readily dispersed by mechanically mixing the same in finely ground form with the poultry feedstuff, or with an intermediate formulation (premix) that is subsequently blended with other components to prepare the final poultry feedstuff that is fed to the poultry. Typical components of poultry feedstuff include molasses, fermentation residues, corn meal, ground and rolled oats, wheat shorts and middlings, alfalfa, clover and meat scraps, together with mineral supplements such as bone meal, calcium carbonate and vitamins.

When the compound according to the present invention is used as an additive to the feed, it is typically incorporated into a "premix." The premix contains the active agent or agents as well as physiologically acceptable carriers and feedstuffs. The premix is relatively concentrated and is adapted to be diluted with other carriers, vitamin and mineral supplements, and feedstuffs to form the final animal feed.

Premixes which are intermediate in concentration of active agent between a first premix and the final animal feed are sometimes employed in the industry and can be used in implementing the present invention. When employing the present compound as sole active agent, a premix desirably contains the agent at a concentration of from 0.1 to 50.0% by weight. Preferred premixes will generally contain the present compound at a concentration of from 0.5 to 25.0%, by weight. The identity of the other components of the premix and ultimate animal feed is not critical. In final

feeds, the concentration of the active agent is not critical and will depend on various factors known to those skilled in the art. Such factors include the relative potency of the particular active agent and the severity of the coccidial challenge. In general, a final feed employing compound of the present invention as the sole anticoccidial will contain from about 0.0005 to about 0.05% by weight of said compound, preferably from about 0.0005 to about 0.005%.

Compositions containing a compound of formula I may also be prepared in powder or liquid concentrate form. In accordance with standard veterinary formulation practice, conventional water soluble excipients, such as lactose or

sucrose, may be incorporated in the powders to improve their physical properties. Thus one embodiment of suitable powders of this invention comprises 50 to 100% w/w, and for example 60 to 80% w/w of the compound and 0 to 50% w/w and for example 20 to 40% w/w of conventional veterinary excipients. These powders may either be added to animal feedstuff, for example by way of an intermediate premix, or diluted in animal drinking water.

Liquid concentrates of this invention suitably contain a water-soluble compound combination and may optionally include a veterinarily acceptable water miscible solvent, for example polyethylene glycol, propylene glycol, glycerol, glycerol formal or such a solvent mixed with up to 30% v/v of ethanol. The liquid concentrates may be administered to the drinking water of animals, particularly poultry.

The present invention contemplates using a compound of formula (I) as sole anticoccidial agent as well as in combination with one or more other anticoccidial agents. Suitable anticoccidials for combination use include, but are not limited to, amprolium, ethopabate, clopidol, meticlorpindol, decoquinate, dinitolmide, halofuginone, lasalocid, maduramicin, monensin, narasin, nicarbazin, chlortetracycline, oxytetracycline, robenidine, salinomycin, semduramicin, and diclazuril. When used in combination with one or more other anticoccidial agent, the compound of formula (I) may be administered at or lower than the effective doses when used alone; for example, the final feed may contain about 0.0001 to about 0.02% by weight, or preferably from about 0.0005 to about 0.005% of a compound of formula (I). Similarly, the second anticoccidial agent in the combination may be used in an amount at or lower than those commonly used as a sole anticoccidial. The combination may be formulated into medicament for poultry use as described previously.

The formulated medicament may contain, in addition to anticoccidial agent(s) other therapeutic or nutritional agents commonly administered to poultry in the feed or drinking water; such other agents may be, for example, parasiticides, antibacterials, and growth promoters.

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The compounds of the invention are prepared by the following reaction scheme(s). All substituents are as defined above unless indicated otherwise.

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SCHEME 1:

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SCHEME 2:

5 SCHEME 3:

SCHEME 4:

5 SCHEME 5:

Pyrimidyl imidazopyrimidines of formula (IA1) and (IA2)may be prepared according to the procedure shown in Scheme 6 below.

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$$R^{21}$$
 R^{21}
 R^{21}
 R^{22}
(IA1)

SCHEME 6:

Pyrimidyl indazoles of formula (IB) may be prepared according to the procedure shown in Scheme 7 below.

$$R^{21}$$
 R^{21}
 R^{22}
 R^{22}
(IB)

SCHEME 7:

Pyrimidyl benzimidazoles of formula (ID) may be prepared according to the procedure shown in Scheme 8 below.

SCHEME 8:

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The following examples illustrate the preparation of some of the compounds of the invention and are not to be construed as limiting the invention disclosed herein.

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INTERMEDIATE COMPOUND 2:

2-Aminopicoline (100g, 924.7mmol) was suspended in methylene chloride (1000mL), cooled to 0°C and treated dropwise with acetic anhydride (94mL, 1000mmol) over a period of 20min., followed by addition of triethyl amine (101g, 1000mmol). The resulting homogeneous solution was warmed up to room temperature and then concentrated to dryness under reduced pressure. The resulting residue was taken up in ethyl acetate (500mL) and water (100mL), and the pH was then adjusted to 6.0 with 2N HCl or NaOH. The organic layer was separated, dried over anhydrous sodium sulfate, filtered, and evaporated. Recrystallization of the residue from ethyl acetate/hexane gave INTERMEDIATE COMPOUND 2 (101g).

INTERMEDIATE COMPOUND 3:

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The amide INTERMEDIATE COMPOUND 2 (62.9g, 419mmol) was dissolved in water (650mL) by warming to 60°. Potassium permanganate (30.6g) was added and the stirred mixture was heated to 75°C. Additional KMnO₄ (30.6g) was added, and the mixture was heated to reflux. After 3h. of reflux, the mixture was cooled to 75°C and additional KMnO₄ (70.2g) was added cautiously in small portions and refluxed for 15h. The mixture was cooled to room temperature, filtered over

celite and extracted with diethyl ether. The aqueous layer was neutralized with 2N HCl to pH 7.0 and evaporated to yield 86g of INTERMEDIATE COMPOUND 3 which was used in the preparation of INTERMEDIATE COMPOUND 4 below without further purification.

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INTERMEDIATE COMPOUND 4:

The acid INTERMEDIATE COMPOUND 3 (10.0g, 55.6mmol) was suspended in absolute ethanol (300mL) at room temperature, HCl gas was bubbled for 10 minutes and then refluxed for 6h. The ethanol was removed under reduced pressure, the resulting viscous liquid was neutralized with std. sodium bicarbonate and the mixture was extracted with ethyl acetate. The organic phase was dried over anhydrous sodium sulfate, filtered and evaporated to yield the ester INTERMEDIATE COMPOUND 4 (2.42g).

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INTERMEDIATE COMPOUND 5:

To a solution of the ester INTERMEDIATE COMPOUND 4 (48.2g, 290mmol) in anhydrous tetrahydrofuran (480mL) at -30°C, lithium aluminum hydride (1.0M in THF, 580mL, 580mmol) was added dropwise. The resulting solution was warmed to 0°C and then refluxed for 1h. The resulting solution was cooled to room temperature, quenched with water (15.4mL) followed by addition of NaOH (5N, 15.4mL) and filtration. The filtrate was evaporated and triturated with diethyl ether to yield the alcohol INTERMEDIATE COMPOUND 5 (25.4g).

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Alternatively, the alcohol INTERMEDIATE COMPOUND 5 could be made from 2-chloro isonicotinic acid by the following sequence of reactions: i) reduction with diborane to alcohol, ii) conversion to tetrazolopyridine with ammonium azide and iii) reduction of the tetrazolopyridine to 2-amino pyridine with zinc in acetic acid or tin dichloride.

INTERMEDIATE COMPOUND 6:

To 2-mercapto-4-methylpyrimidine.HCl (20g, 123mmol) in toluene (300mL) at room temperature under argon, diisopropylethylamine (34.6mL, 184.5mmol) and N,N-dimethylformamide dimethyl acetal (40mL, 301mmol) were added, refluxed for 4h., cooled to room temperature and then concentrated under reduced pressure. The resulting viscous liquid was dissolved in diethyl ether (200mL), diluted with water (50mL) and the pH adjusted to 5.0 with sodium bisulfate (aq. sat.). The organic phase was dried over anhydrous sodium sulfate and concentrated to yield INTERMEDIATE COMPOUND 6 (15.3g) as a light brown oil.

20 INTERMEDIATE COMPOUND 7:

To a solution of INTERMEDIATE COMPOUND 6 (6.3g, 45.0mmol) in THF (100mL, anhydrous) at -78°C under argon, lithium diisopropyl

amide (2.0M in THF, 27.0mL, 54.0mmol) was added dropwise. The resulting solution was stirred for 1hr at -78°C and then treated dropwise with a solution of methyl 4-fluorobenzoate (6.4g, 49.5mmol) in THF (20mL, anhydrous). The mixture was stirred for 2h. at -78°, and then warmed up to room temperature. The resulting solution was quenched with ammonium chloride (aq. sat.) and extracted with ethyl acetate. The organic phase was concentrated and purified by flash column chromatography (silica, 15:85 = EtOAc:Hexane) to yield the ketone INTERMEDIATE COMPOUND 7 (7.58g).

10 INTERMEDIATE COMPOUND 8:

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Tetrabutylammonium tribromide (52.3g, 108mmol) was added to the ketone INTERMEDIATE COMPOUND 7 (28.5g, 108mmol) suspended in carbon tetrachloride (325mL) at room temperature. After 15 minutes, methylene chloride (650mL) was added. The resulting solution was stirred for 4 hours at room temperature. The reaction mixture was quenched with sodium bicarbonate (250mL, sat., aq.) and extracted with ethyl acetate. The organic layer was dried over anhydrous sodium sulfate, filtered and concentrated to a brown oil INTERMEDIATE COMPOUND 8 (78g) which was used in the next step to prepare COMPOUND 9 without further purification.

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INTERMEDIATE COMPOUND 9:

To a solution of the crude bromide INTERMEDIATE COMPOUND 8 (5.45g, 16.0mmol) in absolute ethanol (30mL) at room temperature, the alcohol INTERMEDIATE COMPOUND 5 (894mg, 7.2mmol) dissolved in absolute ethanol (20mL, anhydrous) was added dropwise. The combined solution was heated to 60°C overnight under argon. The resulting solution was diluted with sodium bicarbonate (sat., aq.) and extracted with ethyl acetate. The organic phase was dried over anhydrous sodium sulfate, filtered, concentrated, and purified by flash silica column chromatography (60:40 EtOAc:Hexane first, then 100%EtOAc) to yield the imidazopyridine INTERMEDIATE COMPOUND 9 (806mg).

INTERMEDIATE COMPOUND 10:

15 The imidazopyridine INTERMEDIATE COMPOUND 9 (322mg, 0.88mmol) in methanol (25mL) at room temperature was treated dropwise with a solution of oxone (1082mg, 1.76mmol) in water (10mL). The resulting mixture was stirred at room temperature overnight and extracted with ethyl acetate. The organic phase was dried over anhydrous sodium sulfate, filtered, concentrated, and purified by flash column chromatography (silica, 55:45 EtOAc:Hexane) to yield the sulfone INTERMEDIATE COMPOUND 10 (301mg).

COMPOUND 14 (EXAMPLE A04):

METHOD I:

Step A:

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5 COMPOUND 11 (EXAMPLE A01):

A suspension of the sulfone INTERMEDIATE COMPOUND 10 (300mg, 0.75mmol) in (S)-(-)- alpha-methylbenzylamine (6.0mL) was heated to 60° C for 4h while stirring under an atmosphere of argon. The resulting solution was cooled to room temperature, acidified with citric acid (5%, aq.) to pH = 4.5 and extracted with ethyl acetate. The organic phase was dried over anhydrous sodium sulfate, filtered, concentrated, and purified by flash column chromatography (5:95 10% NH₄OH in MeOH: CH₂Cl₂) to yield the amine **EXAMPLE A01** (307mg).

15 Step B: COMPOUND 12 (EXAMPLE A02):

To a solution of **EXAMPLE A01** (75mg, 0.17mmol) in toluene (3.0mL) at 0°C under argon, 1,8-diazabicyclo[5.4.0]undec-7-ene (0.062mL, 0.40mmol) and diphenylphosphoryl azide (0.088mL, 0.40mmol) were added. The resulting solution was stirred at room temperature overnight, concentrated, and purified by prep silica gel TLC (50:50 EtOAc:Hexane) to give the azide **EXAMPLE A02** (47mg).

Step C:

10 COMPOUND 13 (EXAMPLE A03):

To a solution of the azide EXAMPLE A02 (45mg, 0.10mmol) in THF (1.5mL) at room temperature, triphenylphosphine (65mg, 0.25mmol) and water (1.5mL) were added and stirred at room temperature overnight. The resulting solution was diluted with water, extracted with ethyl acetate, and the organic phase was concentrated and purified by prep silica gel TLC (5:95 = 10% NH₄OH in MeOH: CH₂Cl₂) to yield the amine EXAMPLE A03 (29mg).

Step D:

20 EXAMPLE A04:

To the solution of the amine **EXAMPLE A03** (29mg, 0.10mmol) in methanol (1.0mL) at room temperature under argon, acetic acid (glacial, 0.033mL), formaldehyde (36~38% in water, 0.033mL) and sodium cyanoborohydride (1.0M in THF, 0.52mL, 0.52mmol) were added and stirred at room temperature overnight. The resulting solution was concentrated and purified by prep silica gel TLC (10:90 10% NH₄OH in MeOH: CH₂Cl₂) to yield the dimethyl amine **EXAMPLE A04** (26mg).

10 INTERMEDIATE COMPOUND 18:

METHOD II:

INTERMEDIATE COMPOUND 15:

15 Step A:

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Treatment of INTERMEDIATE COMPOUND 10 with neopentyl amine following the procedure described in Method I, Step A gave INTERMEDIATE COMPOUND 15.

5 Step B:

To a stirred solution of INTERMEDIATE COMPOUND 15 (937mg, 2.31mmol) in chloroform (15mL) at -10°C, was added triethyl amine (0.64mL, 4.62mmol) followed by methane sulfonyl chloride (0.197mL). After 4h., the resulting mesylate COMPOUND 16 (EXAMPLE A06) was treated with dimethylamine (2M in THF, 5mL) and stirring continued overnight at room temperature. The following day, the solution was evaporated and purified by flash column chromatography (silica, 0.6% NH4OH, 5.4% methanol, 94% methylene chloride) to yield INTERMEDIATE COMPOUND 18 (720mg).

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COMPOUND 20 (EXAMPLE A20):

In a pressure vessel, the sulfone INTERMEDIATE COMPOUND 10

20 (1.1g) was suspended in tetrahydrofuran (65mL) saturated with ammonia at -20°C. The tube was closed, warmed up to room temperature and stirred for two days. The vessel was cooled to -35°C, opened, warmed up to room temperature, and then evaporated under reduced pressure. The resulting residue was purified by flash

column chromatography (silica, 9% methanol with 1% ammonium hydroxide, 90% methylene chloride) to yield the amine COMPOUND 19 (EXAMPLE A09) (895mg). Treatment of EXAMPLE A09 (729mg, 2.17mmol) in methylene chloride (10mL) sequentially with triethylamine (0.453, 3.26mmol), methane sulfonyl chloride (0.185mL, 2.39mmol) followed by the treatment of the mesylate with a solution of 2M dimethylamine in tetrahydrofuran as shown in Method II, Step B gave EXAMPLE A20 (215mg).

10 METHOD III COMPOUND 17 (EXAMPLE A07):

15 Step A:

To a solution of **EXAMPLE A01** (150mg, 0.34mmol) in methylene chloride (15mL), manganese dioxide (300mg) was added and stirred for 6h. Filtration over celite and purification by prep TLC (silica, 0.5% NH4OH, 4.5% methanol, 95% methylene chloride) gave the aldehyde **COMPOIUND 17** (**EXAMPLE A07**)

20 (122mg).

Step B:

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COMPOUND 14 (EXAMPLE A04):

To the aldehyde **EXAMPLE A07** (30mg, 0.074mmol) in methylene chloride (1mL) was added dimethyl amine (2M in THF, 0.056mL, 0.117mmol), diisopropylethylamine (0.042mL, 0.222mmol), sodium tiracetoxyborohydride (31.1mg, 0.148mmol) and stirred for 4h. The resulting solution was concentrated and purified by prep silica gel TLC (10:90 10% NH₄OH in MeOH: CH₂Cl₂) to yield the dimethyl amine **EXAMPLE A04** (21mg).

COMPOUND II (EXAMPLE 1):

EXAMPLE 1 was prepared under conditions similar to those used for the synthesis of INTERMEDIATE COMPOUND 15. The key cyclization reaction to form the imidazopyridine ring required the use of 2-amino-3-benzyloxypyridine

(II)

(2.5 equivalents) in isopropanol solvent at a concentration of 0.2 M, heated at 90°C for 14h. The resulting mixture was then concentrated *in vacuo* and purified by flash chromatography (Biotage 40S, SiO₂, 20% EtOAc-hexane) to provide the imidazopyridine cyclization product. This intermediate was elaborated into **EXAMPLE 1** using methodology displayed in **Schemes 2-4** and was characterized by ¹H NMR, HPLC and mass spectrometry (m/z: 482 (M¹+1)).

EXAMPLES 2-32:

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The following imidazopyridines were prepared under conditions 10 similar to those displayed in Schemes 1-5. The 2,4-difluorophenyl moiety of **EXAMPLES 2-6** was introduced by the substitution of methyl 2,4-difluorobenzoate in place of methyl 4-fluorobenzoate shown in Scheme 2. The 3trifluoromethylphenyl moiety of EXAMPLES 7-11 was introduced by the substitution of methyl 3-trifluoromethylbenzoate in place of methyl 4-fluorobenzoate 15 also shown in Scheme 2. The 2-chloro-4-fluorophenyl moiety of EXAMPLES 12-14 was introduced by the substitution of methyl 2-chloro-4-fluorobenzoate in place of methyl 4-fluorobenzoate also shown in Scheme 2. The 2-chlorophenyl moiety of **EXAMPLES 15** and 16 was introduced by the substitution of methyl 2chlorobenzoate in place of methyl 4-fluorobenzoate also shown in Scheme 2. The 4chlorophenyl moiety of EXAMPLES 17 and 18 was introduced by the substitution of 20 methyl 4-chlorobenzoate in place of methyl 4-fluorobenzoate also shown in Scheme 2. The 3,4-dichlorophenyl moiety of EXAMPLES 19 and 20 was introduced by the substitution of methyl 3,4-dichlorobenzoate in place of methyl 4-fluorobenzoate also shown in Scheme 2. The 2,3-dichlorophenyl moiety of EXAMPLES 21 and 22 was introduced by the substitution of methyl 2,3-dichlorobenzoate in place of methyl 4-25 fluorobenzoate also shown in Scheme 2. The cyclohexylamine moiety in **EXAMPLES 4** and 5 was introduced by the substitution of cyclohexylamine in place of neopentylamine shown in Scheme 4. The alcohol moiety in EXAMPLE 27 was introduced by the substitution of (R)-phenyl glycinol in place of neopentylamine 30 shown in Scheme 4. The hydroxyneopentylamine moiety in EXAMPLES 24-26 was introduced by the substitution of 2,2-dimethyl-3-amino-1-propanol in place of neopentylamine displayed in Scheme 4. The sulfonamide moiety in EXAMPLE 26 was introduced by treatment with methanesulfonyl chloride prior to the introduction of the 2,2-dimethyl-3-amino-1-propanol subunit. The methyl ether moiety in 35 **EXAMPLE 6** was introduced by the substitution of sodium methoxide in place of

dimethylamine displayed in Scheme 4. The methyl sulfone moiety in EXAMPLE 31 was introduced by the substitution of sodium thiolate in place of dimethylamine shown in Scheme 4 to provide the methyl sulfide intermediate. This methyl sulfide was then oxidized with 2 equivalents of oxone in 2:1 methanol-water to provide the methyl sulfone in EXAMPLE 31. The dimethyl phosphonate moiety in EXAMPLE 5 30 was introduced by the substitution of sodium dimethylphosphite in place of dimethylamine described in Scheme 4. The morpholine moiety in EXAMPLE 7 was introduced by the substitution of morpholine in place of dimethylamine described in Scheme 4. The dimethylaminoethylpiperazine moiety in EXAMPLE 8 was 10 introduced by the substitution of N-(2-(N,N-dimethylamino)ethyl)piperazine in place of dimethylamine described in Scheme 4. The isopropylpiperazine moiety in **EXAMPLE 9** was introduced by the substitution of N-isopropylpiperazine in place of dimethylamine shown in Scheme 4. The methylamine moiety in EXAMPLE 25 was introduced by the substitution of methylamine (2M in THF) in place of dimethylamine 15 shown in Scheme 4. The sulfonamide moiety in Example 28 was introduced by treating the analogous neopentyl derivative of EXAMPLE A03 (COMPOUND 13) shown in Scheme 3 with methanesulfonyl chloride. This sulfonamide was subsequently alkylated with KHMDS/MeI to provide EXAMPLE 29. EXAMPLES 23-24 (Z=H) and 27 (Z=CH₃) were prepared under conditions where 2-amino-4-20 hydroxymethylpyridine (INTERMEDIATE COMPOUND 5) was substituted by 2aminopyridine and 2-amino-4-picoline respectively as shown in Scheme 2. The aldehydes in EXAMPLES 13 and 32 were prepared by oxidation of EXAMPLE 12 and INTERMEDIATE COMPOUND 15 (Scheme 4) using Dess-Martin periodinane in methylene chloride in a similar manner shown in Scheme 5. With the 25 exception of EXAMPLE 16 (1H NMR only), the following imidazopyridines were characterized by ¹H NMR, HPLC and mass spectrometry.

The following **TABLE 1** of **EXAMPLES 2-32** refer to the following general chemical structure:

TABLE 1

TABLE 1				
EX.	Ar Group	R Group	Z Group	MS (m/z)
2	2,4-Difluorophenyl	NHCH ₂ C(CH ₃) ₃	СН₂ОН	424 (M ⁺ +1)
3	2,4-Difluorophenyl	NHCH ₂ C(CH ₃) ₃	CH ₂ N(CH ₃) ₂	451 (M ⁺ +1)
4	2,4-Difluorophenyl	HN	CH₂OH	436 (M ⁺ +1)
5	2,4-Difluorophenyl	HN	CH ₂ N(CH ₃) ₂	463 (M ⁺ +1)
6	2,4-Difluorophenyl	NHCH ₂ C(CH ₃) ₃	CH₂OCH₃	438 (M ⁺ +1)
7	3- Trifluoromethylphenyl	NHCH₂C(CH₃)₃	CH ₂ -NO	525 (M ⁺ +1)
8	3- Trifluoromethylphenyl	NHCH ₂ C(CH ₃) ₃	CH ₂ -N_N-NMe ₂	595 (M ⁺ +1)
9	3- Trifluoromethylphenyl	NHCH ₂ C(CH ₃) ₃	CH ₂ -N N-	566 (M ⁺ +1)
10	3- Trifluoromethylphenyl	NHCH₂C(CH₃)₃	СН₂ОН	458 (M ⁺ +1)
11	3- Trifluoromethylphenyl	NHCH ₂ C(CH ₃) ₃	CH ₂ N(CH ₃) ₂	483 (M ⁺ +1)
12	2-Chloro-4- fluorophenyl	NHCH ₂ C(CH ₃) ₃	СН₂ОН	442 (M ⁺ +1)
13	2-Chloro-4- fluorophenyl	NHCH₂C(CH₃)₃	СНО .	438 (M ⁺ +1)
14	2-Chloro-4- fluorophenyl	NHCH₂C(CH₃)₃	CH ₂ N(CH ₃) ₂	467 (M ⁺ +1)
15	2-Chlorophenyl	NHCH₂C(CH₃)₃	СН₂ОН	422 (M ⁺ +1)
16	2-Chlorophenyl	NHCH ₂ C(CH ₃) ₃	CH ₂ N(CH ₃) ₂	
17	4-Chlorophenyl	NHCH ₂ C(CH ₃) ₃	СН₂ОН	422 (M ⁺ +1)
18	4-Chlorophenyl	NHCH ₂ C(CH ₃) ₃	CH ₂ N(CH ₃) ₂	450 (M ⁺ +1)
19	3,4-Dichlorophenyl	NHCH₂C(CH₃)₃	СН₂ОН	457 (M ⁺ +1)
20	3,4-Dichlorophenyl	NHCH₂C(CH₃)₃	CH ₂ N(CH ₃) ₂	484 (M ⁺ +1)
21	2,3-Dichlorophenyl	NHCH ₂ C(CH ₃) ₃	СН₂ОН	457 (M ⁺ +1)
22	2,3-Dichlorophenyl	NHCH₂C(CH₃)₃	CH₂N(CH₃)₂	484 (M ⁺ +1)
23	4-Fluorophenyl	NHCH₂C(CH₃)₃	н	376 (M ⁺ +1)

EX.	Ar Group	R Group	Z Group	MS (m/z)
24	4-Fluorophenyl	NHCH ₂ C(CH ₃) ₂ CH ₂ OH	Н	392 (M ⁺ +1)
25	4-Fluorophenyl	NHCH₂C(CH₃)₂CH₂OH	CH₂NHCH₃	435 (M ⁺ +1)
26	4-Fluorophenyl	NHCH₂C(CH₃)₂CH₂OH	CH ₂ N(CH ₃)SO ₂ CH	513 (M ⁺ +1)
Ŀ			3	
27	4-Fluorophenyl	HN	CH ₃	440 (M ⁺ +1)
		(R)		·
28	4-Fluorophenyl	NHCH₂C(CH₃)₃	CH2NHSO2CH3	483 (M ⁺ +1)
29	4-Fluorophenyl	NHCH ₂ C(CH ₃) ₃	CH ₂ N(CH ₃)SO ₂ CH	497 (M ⁺ +1)
			3	
30	4-Fluorophenyl	NHCH₂C(CH₃)₃	CH ₂ PO(OMe) ₂	498 (M ⁺ +1)
31	4-Fluorophenyl	NHCH ₂ C(CH ₃) ₃	CH ₂ SO ₂ CH ₃	468 (M ⁺ +1)
32	4-Fluorophenyl	NHCH₂C(CH₃)₃	СНО	404 (M ⁺ +1)

EXAMPLE 33A (COMPOUND III) and EXAMPLE 33B (COMPOUND IV):

(III)

(IV)

Compounds (III) and (IV) were prepared from INTERMEDIATE

5 COMPOUND 7 as shown in Schemes 2 and 6. Thus, the methylsulfide
INTERMEDIATE COMPOUND 7 (8g, 30.7mmol) was diluted into 2:1 MeOHH₂O (700mL), oxone (38g, 61.4mmol) added, and the suspension stirred at 23°C for
15h. The resulting reaction mixture was concentrated in vacuo, and the residue
purified by flash column chromatography (Biotage 40M, SiO₂, 50% EtOAc-hexane)
to provide the sulfone intermediate (6.8g). This material (6.8g, 23.2mmol) was
diluted into dichloroethane (100mL) and neopentylamine (6.1g, 69.5mmol) added.

The resulting reaction mixture was heated at 50°C for 15h., cooled, partitioned between aqueous sodium bicarbonate and methylene chloride, the organic phase dried with anhydrous sodium sulfate, and concentrated *in vacuo*. The residue was purified by flash column chromatography (Biotage 40M, SiO₂, 15% EtOAchexane) to provide the aminopyrimidine intermediate (2g). This material (1.9g, 6.45mmol) was diluted into 2:1 methylene chloride-CCl₄ (60mL) and treated with tetrabutylammonium tribromide (3.4g, 7.1mmol) added. The reaction mixture was maintained at 23°C for 30min., partitioned between aqueous sodium bicarbonate and methylene chloride, the organic phase dried with anhydrous sodium sulfate, and concentrated *in vacuo*.

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The resulting residue was purified by flash column chromatography (Biotage 40M, SiO₂, 5-20% EtOAc-hexane) to provide the bromide intermediate (2.2g). This material (200mg, 0.53mmol) was diluted into NMP (0.53mL) and treated with 2-aminopyrimidine (505mg, 5.3mmol). The resulting reaction mixture was maintained at 135°C for 4h., cooled, and purified by flash column chromatography

(Biotage 40M, SiO₂, 20% EtOAc-hexane) to provide a mixture of two regioisomeric products. This mixture was separated by preparative thin layer chromatography (3 X 1500u, SiO₂, 2% methanol-chloroform) to provide (III) and (IV) which were each characterized by ¹H NMR, HPLC and mass spectrometry (m/z: 377 (M⁺+1)) for (III) and (m/z: 377 (M⁺+1)) for (IV).

EXAMPLE 34 (COMPOUND V).

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(V)

Compound (V) was prepared under conditions similar to those used for the synthesis of INTERMEDIATE COMPOUND 15. The key cyclization reaction to form the imidazoisoquinoline ring required the use of isoquinolin-3-amine (2.5 equivalents) in isopropanol solvent at a concentration of 0.2M, heated at 90°C for 14h. The mixture was then concentrated in vacuo and purified by flash chromatography (Biotage 40S, SiO₂, 20% EtOAc-hexane) to provide the imidazoisoquinoline cyclization product. This intermediate was elaborated into (V) using methodology displayed in Schemes 2-4 and was characterized by ¹H NMR, HPLC and mass spectrometry (m/z: 426 (M⁺+1)).

EXAMPLE 35 (COMPOUND VI):

(VI)

for the synthesis of INTERMEDIATE COMPOUND 15. The key cyclization reaction to form the imidazoisoquinoline ring required the use of 1-aminoisoquinoline (2.5 equivalents) in isopropanol solvent at a concentration of 0.2M, heated at 90°C for 14h. The resulting mixture was then concentrated *in vacuo* and purified by flash chromatography (Biotage 40S, SiO₂, 20% EtOAc-hexane) to provide the imidazoisoquinoline cyclization product. This intermediate was elaborated into (VI) using methodology displayed in Schemes 2-4 and was characterized by ¹H NMR, HPLC and mass spectrometry (m/z: 426 (M⁺+1)).

15 EXAMPLE 36 (COMPOUND VII):

(VII)

Compound (VII) was prepared under conditions similar to those used for the synthesis of INTERMEDIATE COMPOUND 15. The key cyclization

5 reaction to form the imidazotriazine ring required the use of 3-amino-1,2,4-triazine (2.5 equivalents) in isopropanol solvent at a concentration of 0.2M, heated at 90°C for 14h. The resulting mixture was then concentrated *in vacuo* and purified by flash chromatography (Biotage 40M, SiO₂, 70% EtOAc-hexane) to provide the imidazotriazine cyclization product in 38% yield. This intermediate was elaborated into (VII) using methodology displayed in Schemes 2-4 and was characterized by ¹H NMR, HPLC and mass spectrometry (m/z: 378 (M⁺+1)).

EXAMPLE 37 (COMPOUND VIII):

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(VIII)

Compound (VIII) was prepared under conditions similar to those used for the synthesis of INTERMEDIATE COMPOUND 15. The key cyclization reaction to form the imidazobenzimidazole ring required the use of 2-amino-1-methylbenzimidazole (2.5 equivalents) in isopropanol solvent at a concentration of 0.2M, heated at 90°C for 14h. The resulting mixture was then concentrated *in vacuo* and purified by flash chromatography (Biotage 40M, SiO₂, 70% EtOAc-hexane) to provide the intermediate hydrated pre-cyclization product. This intermediate was dehydrated with Burgess Reagent (5 equivalents) in dioxane at 90°C for 12h to form the imidazobenzimidazole cyclization product (30% yield) which was elaborated into

(VIII) using methodology displayed in Schemes 2-4 and was characterized by ¹H NMR, HPLC and mass spectrometry (m/z: 429 (M⁺+1)).

EXAMPLE 38 (COMPOUND IX):

(IX)

Compound (IX) was prepared under conditions similar to those used for the synthesis of INTERMEDIATE COMPOUND 15. The key cyclization reaction to form the imidazothiazole ring required the use of 2-aminothiazole (3 equivalents) in isopropanol solvent at a concentration of 0.2M, heated at 90°C for 14h. The mixture was then concentrated *in vacuo* and purified by flash chromatography (Biotage 40M, SiO₂, 25% EtOAc-hexane) to provide the imidazotriazine cyclization product in 41% yield. This intermediate was elaborated into (IX) using methodology displayed in Schemes 2-4 and was characterized by ¹H NMR, HPLC and mass spectrometry (m/z: 382 (M⁺+1)).

EXAMPLES 39-64:

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The following imidazothiazoles were prepared under conditions similar to those described in **EXAMPLE 38** for the synthesis of **(IX)**. The 2,4-difluorophenyl moiety of **EXAMPLE 39** was introduced by the substitution of methyl 2,4-difluorobenzoate in place of methyl 4-fluorobenzoate shown in **Scheme 2**. The 3-trifluoromethylphenyl moiety of **EXAMPLES 40-42** was introduced by the substitution of methyl 3-trifluoromethylbenzoate in place of methyl 4-fluorobenzoate shown in **Scheme 2**. The R-Groups in **EXAMPLES 41-64** were introduced by the

substitution of the respective amines in place of neopentylamine shown in Scheme 4. The following imidazothiazoles were characterized by ¹H NMR, HPLC and mass spectrometry.

The following **TABLE 2** of **EXAMPLES 39-64** refer to the following general chemical structure:

TABLE 2

Yayı	A - C	D.C.	
EX.	Ar Group	R Group	MS (m/z)
39	2,4-Difluorophenyl	NHCH ₂ C(CH ₃) ₃	400 (M ⁺ +1)
40 .	3-Trifluoromethylphenyl	NHCH ₂ C(CH ₃) ₃	432 (M ⁺ +1)
41	3-Trifluoromethylphenyl	(S) NH	466 (M ⁺ +1)
42	3-Trifluoromethylphenyl	NH(CH ₂) ₃ OCH ₃	434 (M ⁺ +1)
43	4-Fluorophenyl	NH	436 (M ⁺ +1)
		CI	APP CONTRACTOR
44	4-Fluorophenyl	F ₃ C NH	470 (M ⁺ +1)
45	4-Fluorophenyl	HN OH (R,S)	446 (M ⁺ +1)
46	4-Fluorophenyl	NHCH₂C(CH₃)₂CH₂OH	398 (M ⁺ +1)
47	4-Fluorophenyl	(S) NH	466 (M ⁺ +1)

EX.	Ar Group	R Group	MS (m/z)
48	4-Fluorophenyl	(S) NH	466 (M ⁺ +1)
49	4-Fluorophenyl	(S) NH	416 (M ⁺ +1)
50	4-Fluorophenyl	NH(CH ₂) ₃ OCH ₃	384 (M ⁺ +1)
51	4-Fluorophenyl	NH	380 (M ⁺ +1)
52	4-Fluorophenyl	HN	394 (M ⁺ +1)
53	4-Fluorophenyl	H₃C NH	416 (M ⁺ +1)
54	4-Fluorophenyl	HN OH (R)	432 (M ⁺ +1)
55	4-Fluorophenyl	F ₃ C S	502 (M ⁺ +1)
56	4-Fluorophenyl	NH CO ₂ H	446 (M ⁺ +1)
57	4-Fluorophenyl	HN OH (S)	460 (M ⁺ +1)
58	4-Fluorophenyl	HN OH OH (R)	460 (M ⁺ +1)
59	4-Fluorophenyl	OH HN,,, O OH (s,s)	476 (M ⁺ +1)

EX.	Ar Group	R Group	MS (m/z)
60	4-Fluorophenyl	MeS	448 (M ⁺ +1)
61	4-Fluorophenyl	HN OMe (R)	446 (M ⁺ +1)
62	4-Fluorophenyl	HN (R,S)	412 (M ⁺ +1)
63	4-Fluorophenyi	HN (R,S)	410 (M ⁺ +1)
64	4-Fluorophenyl	HN	396 (M ⁺ +1)

INTERMEDIATE COMPOUND X:

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To a solution of Me(MeO)NH-HCl (4.9g, 50.7mmol), EDCI (2.1g, 11.2mmol) and DIEA (10.6mL, 60.9mmol) in 1:1 DMF-CH₂Cl₂ (75mL) at 0°C was added 3-nitro-4-(hydroxymethyl)benzoic acid (2g, 10.1mmol) in 1:1 DMF-CH₂Cl₂ (50mL). The resulting reaction mixture was warmed to 23°C, maintained 15h., partitioned between NH₄Cl_(aq) and CH₂Cl₂, the organic phase washed with NaHCO_{3(aq)}, then dried over anhydrous sodium sulfate and concentrated *in vacuo*. The crude product (1.4g, 5.8mmol) was then diluted into acetonitrile (40mL) and treated with MnO₂ (2.5g, 29.2mmol). The resulting reaction mixture was maintained at 23°C for 15h., filtered through celite and concentrated *in vacuo*. The crude material was purified by flash column chromatography (SiO₂, acetone-hexane) to provide

880mg of product which was characterized by ¹H NMR, HPLC and mass spectrometry (m/z: 239 (M⁺+1)).

This material (800mg, 3.3mmol) was then diluted into toluene (3.3mL), 4-fluoroaniline (0.35mL, 3.6mmol) was added, and the resulting reaction mixture was heated at 100°C. Concentration *in vacuo* of the reaction mixture provided 900mg (2.7mmol) of crude product which was diluted into triethyl phosphite (3mL) and heated at 150°C for 15h., the excess triethyl phosphite removed by distillation, and the residue purified by flash column chromatography (SiO₂, acetonehexane) to provide 680mg of (X) which was characterized by ¹H NMR, HPLC and mass spectrometry (m/z: 300 (M⁺+1)).

INTERMEDIATE COMPOUND XI:

(XI)

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INTERMEDIATE COMPOUND X (660mg, 2.2mmol) was diluted into glacial acetic acid (15mL) and slowly treated with a solution of bromine (0.11mL, 350mg, 2.2mmol) in glacial acetic acid (10mL) over 3h. The resulting reaction mixture was maintained at 23°C for 15h., poured into ice water, filtered, and the residue purified by flash column chromatography (SiO₂, acetone-hexane) to provide 550mg of product which was characterized by ¹H NMR, HPLC and mass spectrometry (m/z: 378 (M⁺+1)).

This material (25mg, 0.068mmol) was then diluted into DMF (0.5mL), 2-chloro-4-(trimethylstannyl)pyrimidine (38mg, 0.14mmol) was added, followed by Pd₂(dba)₃ (4mg) and P(o-tol)₃ (2.5mg), and the reaction mixture was heated at 100°C. The reaction mixture was concentrated *in vacuo* and the residue purified by preparative thin layer chromatography (SiO₂, 5% MeOH-CH₂Cl₂) to provide 20mg of

(XI) which was characterized by ¹H NMR, HPLC and mass spectrometry (m/z: 412 (M⁺+1)).

EXAMPLE 67 (COMPOUND XII):

(XII)

COMPOUND XI (20mg, 0.049mmol) was diluted into DMSO (0.5mL) and treated with neopentylamine (0.011mL, 0.097mmol). The resulting reaction mixture was maintained at 100°C for 15h., and the reaction mixture partitioned between water and chloroform, the organic phase dried over anhydrous sodium sulfate and concentrated *in vacuo*. The residue was purified by preparative thin layer chromatography (SiO₂, 5% MeOH-CH₂Cl₂) to provide 10mg of COMPOUND XII which was characterized by ¹H NMR, HPLC and mass spectrometry (m/z: 463 (M⁺+1)).

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EXAMPLE 68 (COMPOUND XIII):

(XIII)

COMPOUND XII (45mg, 0.097mmol) was diluted into toluene (1.5mL), cooled to -78°C and treated with DIBAL-H (1 M in toluene, 0.107mL, 0.107mmol). The resulting reaction mixture was maintained at -78°C for 1h., and then quenched with aqueous potassium sodium tartrate (0.060mL), warmed to 23°C, filtered through celite, washed with Et₂O, the solution then dried over anhydrous sodium sulfate and concentrated *in vacuo*. The residue was purified by preparative thin layer chromatography (SiO₂, acetone-hexane) to provide 20mg of COMPOUND XIII which was used directly in EXAMPLE 69 below.

EXAMPLE 69 (COMPOUND XIV).

(XIV)

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COMPOUND XIII (25mg, 0.062mmol) was diluted into THF (1mL) and treated with NaBH₄ (24mg, 0.62mmol). The resulting reaction mixture was maintained at 23°C for 1h., partitioned between aqueous sodium bicarbonate and methylene chloride, dried over anhydrous sodium sulfate and concentrated *in vacuo*.

The residue was purified by preparative thin layer chromatography (SiO₂, 5% MeOH-chloroform) to provide 15mg of **COMPOUND XIV** which was characterized as two isomers by ¹H NMR, HPLC and mass spectrometry (m/z: 406 (M⁺+1)).

5 EXAMPLE 70 (COMPOUND XV):

(XV)

COMPOUND XIII (20mg, 0.050mmol) was diluted into CH₂Cl₂

(1mL) and treated with dimethylamine (0.037mL, 0.074mmol), DIEA (0.030mL, 0.150mmol) and Na(OAc)₃BH (21mg, 0.10mmol). The resulting reaction mixture was maintained at 23°C for 4h., partitioned between aqueous sodium bicarbonate and methylene chloride, dried over anhydrous sodium sulfate and concentrated *in vacuo*. The residue was purified by preparative thin layer chromatography (SiO₂, 5% MeOH-chloroform) to provide 21mg of COMPOUND XV which was characterized by ¹H NMR, HPLC and mass spectrometry (m/z: 433 (M⁺+1)).

EXAMPLES 71-78:

The following indazoles were prepared under conditions similar to
those described in **EXAMPLES 65-70** as shown in **Scheme 7** and were characterized by ¹H NMR, HPLC and mass spectrometry. The **TABLE 3** below for **EXAMPLES**65-70 refer to the following general chemical formula:

TABLE 3

TABLE 3					
EX.	R Group	Z Group	MS (m/z)		
71	NHCH ₂ C(CH ₃) ₃	CON(OMe)Me	481 (M ⁺ +1)		
72	NHCH ₂ C(CH ₃) ₃	СНО	422 (M ⁺ +1)		
73	NHCH2C(CH3)3	CH₂OH	424 (M ⁺ +1)		
74	NHCH ₂ C(CH ₃) ₃	CH ₂ N(CH ₃) ₂	451 (M ⁺ +1)		
75	HN (S)	CON(OMe)Me	515 (M ⁺ +1)		
76	HN (S)	СНО	456 (M ⁺ +1)		
77	HN (S)	СН₂ОН	458 (M ⁺ +1)		
78	HN (S)	CH ₂ N(CH ₃) ₂	485 (M ⁺ +1)		

5 INTERMEDIATE COMPOUND XVI:

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(XVI)

To a solution of Me(MeO)NH-HCl (26.8g, 275mmol), EDCI (10.6g, 54.9mmol) and DIEA (67mL, 384.6mmol) in 1:4 DMF-CH₂Cl₂ (75mL) at 0°C was added 3-nitro-4-aminobenzoic acid (10g, 54.9mmol) in 1:1 DMF-CH₂Cl₂ (50mL). The resulting reaction mixture was warmed to 23°C, maintained 15h., partitioned between NH₄Cl_(aq) and CH₂Cl₂, the organic phase washed with NaHCO_{3(aq)}, then dried over anhydrous sodium sulfate and concentrated *in vacuo*. The crude material was purified by flash column chromatography (SiO₂, acetone-hexane) to provide 8.9g of product which was characterized by ¹H NMR, HPLC and mass spectrometry (m/z: 226 (M⁺+1)).

This material (10g, 44.4mmol) was then diluted into dioxane (100mL), 2-(methylthio)-4-chloropyrimidine (8.6g, 53mmol) was added, followed by cesium carbonate (25.7g, 133mmol), Pd₂(dba)₃ (900mg) and XANTHPHOS (1g), and the resulting reaction mixture was heated at 90°C for 15h. The reaction mixture was partitioned between water and methylene chloride, the organic phase dried over anhydrous sodium sulfate, concentrated *in vacuo* and the solid purified by recrystallization from acetone-hexane (primary) and then ethyl acetate-hexane (secondary) to provide 10.3g of INTERMEDIATE COMPOUND XVI which was characterized by ¹H NMR, HPLC and mass spectrometry (m/z: 350 (M⁺+1)).

INTERMEDIATE COMPOUND XVII:

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(XVII)

INTERMEDIATE COMPOUND XVI (1g, 3.13mmol) was diluted into CH₂Cl₂ (75mL) and treated with Pd-C (300mg), vacuum-purged with hydrogen gas via a balloon, and the resulting reaction mixture maintained at 23°C for 15h under 1atm of hydrogen. The reaction mixture was filtered through celite, and the residue purified by flash column chromatography (SiO2, acetone-hexane) to provide 980mg of product. This material (2.7g, 9.2mmol) was diluted into nitrobenzene (15mL), 2,4difluorobenzaldehyde (1.4g, 10.1mmol) was added, and the resulting reaction mixture was heated at 175°C for 15h. The reaction mixture was loaded directly on silica gel and purified by flash column chromatography (SiO2, acetone-hexane) to provide 1 g of product and 1.6g of intermediate imine. This imine was recycled through the reaction conditions and purified to provide an additional 1.1g of product (total 2.1g of INTERMEDIATE COMPOUND XVII) which was characterized by 'H NMR,

15 HPLC and mass spectrometry (m/z: 442 (M⁺+1)).

EXAMPLE 81 (COMPOUND XVIII):

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(XVIII)

20 INTERMEDIATE COMPOUND XVII (2g, 4.54mmol) was diluted into CH₂Cl₂ (15mL) and methanol (150mL), and treated with a solution of oxone (5.6g, 9.1 mmol) in water (75mL). The resulting reaction mixture was maintained at 23°C for 15h., and the reaction mixture was filtered to remove the precipitate, the filtrate evaporated and then partitioned between aqueous sodium bicarbonate and 25 methylene chloride, the organic phase dried over anhydrous sodium sulfate and

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concentrated *in vacuo*. The residue was purified by flash column chromatography (SiO₂, acetone-hexane) to provide 600mg of sulfone. This material (50mg, 0.106mmol) was diluted into DMSO (1mL) and treated with (S)-(-)-alphamethylbenzylamine (64mg, 0.528mmol). The resulting reaction mixture was maintained at 80°C for 15h., and the reaction mixture partitioned between water and methylene chloride, the organic phase dried over anhydrous sodium sulfate and concentrated *in vacuo*. The residue was purified by preparative thin layer chromatography (SiO₂, acetone-hexane) to provide 48mg of **COMPOUND XVIII** which was characterized by ¹H NMR, HPLC and mass spectrometry (m/z: 515 (M⁺+1)).

EXAMPLE 82 (COMPOUND XIX):

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Representative Procedure from Scheme 8. The starting methoxy methyl amide (60mg, 0.12mmol) was diluted into toluene (1.5mL), cooled to -78°C and treated with DIBAL-H (1 M in toluene, 0.142mL, 0.142mmol). The resulting reaction mixture was maintained at -78°C for 1h., and then quenched with aqueous potassium sodium tartrate (0.022mL), warmed to 23°C, filtered through celite, washed with Et₂O, the solution then dried over anhydrous sodium sulfate and concentrated *in vacuo*. The product (COMPOUND XIX) (56mg) was used directly in EXAMPLE 83 to generate COMPOUND XX below.

(XIX)

25 EXAMPLE 83 (COMPOUND XX).

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(XX)

Representative Procedure from Scheme 8. COMPOUND XIX

5 (25mg, 0.055mmol) was diluted into THF (1mL) and treated with NaBH₄ (21mg, 0.55mmol). The resulting reaction mixture was maintained at 23°C for 1h., partitioned between aqueous sodium bicarbonate and methylene chloride, dried over anhydrous sodium sulfate and concentrated in vacuo. The residue was purified by preparative thin layer chromatography (SiO₂, 5% MeOH-chloroform) to provide 15mg of COMPOUND XX which was characterized by ¹H NMR, HPLC and mass spectrometry (m/z: 456 (M¹+1)).

EXAMPLE 84 (COMPOUND XXI):

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(XXI)

Representative Procedure from Scheme 8. The requisite aldehyde (30mg, 0.074mmol) was diluted into CH₂Cl₂ (1mL) and treated with dimethylamine

(0.056mL, 0.117mmol), DIEA (0.042mL, 0.222mmol) and Na(OAc)₃BH (31mg, 0.15mmol). The resulting reaction mixture was maintained at 23°C for 4h., partitioned between aqueous sodium bicarbonate and ethyl acetate, dried over anhydrous sodium sulfate and concentrated *in vacuo*. The residue was purified by preparative thin layer chromatography (SiO₂, 10% MeOH-chloroform) to provide 21mg of COMPOUND XXI which was characterized by ¹H NMR, HPLC and mass spectrometry (m/z: 433 (M⁺+1)).

EXAMPLES 85-125:

The following benzimidazoles were prepared under conditions similar to those described in **EXAMPLES 79-84** as shown in **Scheme 8** and were characterized by ¹H NMR, HPLC and mass spectrometry. The following **TABLE 4** refers to the following chemical structure:

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TABLE 4

	TADLE 4				
EX	Ar Group	R Group	Z Group	Ms (m/z)	
85	4-Fluorophenyl	NHCH ₂ C(CH ₃) ₃	СН₂ОН	406 (M ⁺ +1)	
86	4-Fluorophenyl	NHCH ₂ C(CH ₃) ₃	CON(OMe)Me	463 (M ⁺ +1)	
87	3-	NHCH ₂ C(CH ₃) ₃	CON(OMe)Me	513 (M ⁺ +1)	
88	Trifluoromethylphenyl 3- Trifluoromethylphenyl	NHCH₂C(CH₃)₃	CH ₂ N(CH ₃) ₂	483 (M ⁺ +1)	
89	2-Chlorophenyi	NHCH ₂ C(CH ₃) ₃	CH ₂ N(CH ₃) ₂	450 (M ⁺ +1)	
90	2-Chloro-4- fluorophenyl	NHCH ₂ C(CH ₃) ₃	CH ₂ N(CH ₃) ₂	467 (M ⁺ +1)	
91	2,4-Difluorophenyl	NHCH ₂ C(CH ₃) ₃	CH ₂ N(CH ₃) ₂	451 (M ⁺ +1)	
92	2,4-Difluorophenyl	NHCH ₂ C(CH ₃) ₃	CH₂OH	424 (M ⁺ +1)	

EX	Ar Group	R Group	Z Group	Ms (m/z)
93	2,4-Difluorophenyl	NHCH2C(CH3)3	CON(OMe)Me	481 (M ⁺ +1)
94	2,4-Difluorophenyl	NHCH₂C(CH₃)₃	CH ₂ -NN-	534 (M ⁺ +1)
95	2,4-Difluorophenyl	NHCH ₂ C(CH ₃) ₃	CH2-N N-NMez	563 (M ⁺ +1)
96	2,4-Difluorophenyl	NHCH ₂ C(CH ₃) ₃	CH ₂ -NNH	504 (M ⁺ +1)
97	2,4-Difluorophenyl	NHCH ₂ C(CH ₃) ₃	CH2-N_O	493 (M ⁺ +1)
98	2,4-Difluorophenyl	NHCH ₂ C(CH ₃) ₃	CH ₂ -NH	532 (M ⁺ +1)
99	2,4-Difluorophenyl	NHCH₂C(CH₃)₃	CH ₂ NH(CH ₂) ₂ OCH ₃	481 (M ⁺ +1)
100	2,4-Difluorophenyl	NHCH₂C(CH₃)₃	CH ₂ NH(CH ₂) ₂ N(C H ₃) ₂	494 (M ⁺ +1)
101	2,4-Difluorophenyl	NH(CH ₂) ₃ OCH ₃	CON(OMe)Me	483 (M ⁺ +1)
102	2,4-Difluorophenyl	H ₃ C NH	CON(OMe)Me	515 (M ⁺ +1)
103	2,4-Difluorophenyl	HN OH (R)	CON(OMe)Me	531 (M ⁺ +1)
104	2,4-Difluorophenyl	NHCH ₂ C(CH ₃) ₂ CH ₂ OH	СН₂ОН	440 (M ⁺ +1)
105	2,4-Difluorophenyl	HN	CON(OMe)Me	495 (M ⁺ +1)
106	2,4-Difluorophenyl	HN (S)	CH ₂ N(CH ₃) ₂	485 (M ⁺ +1)
107	2,4-Difluorophenyl	HN (S)	CH ₂ −NO	527 (M ⁺ +1)
108	2,4-Difluorophenyl	HN (S)	CH ₂ -NNH	538 (M ⁺ +1)

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EX	Ar Group	R Group	Z Group	Ms (m/z)
109	2,4-Difluorophenyl	NHCH ₂ C(CH ₃) ₂ CH	CH ₂ N(CH ₃) ₂	467 (M ⁺ +1)
l		₂ OH		
110	2,4-Difluorophenyl	NH(CH ₂) ₃ OCH ₃	CH2-N_O	495 (M ⁺ +1)
111	2,4-Difluorophenyl	NH(CH ₂) ₃ OCH ₃	CH ₂ N(CH ₃) ₂	453 (M ⁺ +1)
112	2,4-Difluorophenyl	NHCH2C(CH3)2CH	CON(OMe)Me	497 (M ⁺ +1)
		₂ OH		
113	2,4-Difluorophenyl	NH(CH ₂) ₄ OH	CH ₂ N(CH ₃) ₂	453 (M ⁺ +1)
114	2,4-Difluorophenyl	NH	CH ₂ N(CH ₃) ₂	485 (M ⁺ +1)
•		H ₃ C		
115	2,4-Difluorophenyl	HN	CH ₂ N(CH ₃) ₂	501 (M ⁺ +1)
		(R)		
116	2,4-Difluorophenyl	HN	CH ₂ N(CH ₃) ₂	463 (M ⁺ +1)
117	2,4-Difluorophenyl	HN	CH ₂ N(CH ₃) ₂	465 (M ⁺ +1)
118	2,4-Difluorophenyl	HN (S)	CH ₂ N(CH ₃) ₂	515 (M ⁺ +1)
		OMe		
119	2,4-Difluorophenyl	HN (S)	CH ₂ N(CH ₃) ₂	530 (M ⁺ +1)
<u> </u>		NO ₂		
120	2,4-Difluorophenyl	(S)	CH ₂ N(CH ₃) ₂	499 (M ⁺ -15)
		MeO		
121	2,4-Difluorophenyl	HN (S)	CH ₂ N(CH ₃) ₂	564 (M ⁺ +1)
			·	
L		<u>Br</u>	l—————	<u> </u>

EX	Ar Group	R Group	Z Group	Ms (m/z)
122	2,4-Difluorophenyl	HN (S)	CH ₂ N(CH ₃) ₂	510 (M ⁺ +1)
123	2,4-Difluorophenyl	HN (S)	CH ₂ N(CH ₃) ₂	543 (M ⁺ +1)
124	2,4-Difluorophenyl	HN (S)	CH ₂ N(CH ₃) ₂	543 (M ⁺ +1)
125	2,4-Difluorophenyl	NH(CH ₂) ₃ CO ₂ H	CH ₂ N(CH ₃) ₂	467 (M ⁺ +1)

Scheme 10:

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INTERMEDIATE COMPOUND 106:

INTERMEDIATE COMPOUND 106 was prepared by the literature

procedure: J. Med. Chem. 1999, 42 2180-2190.

INTERMEDIATE COMPOUND 107:

INTERMEDIATE COMPOUND 107 was prepared from INTERMEDIATE COMPOUND 106 (10g, 32mmol) using a procedure like that described for the preparation of INTERMEDIATE COMPOUND 8 above in Scheme 2. Yellow oil, (9.91g).

¹H NMR (CDCl₃, 300MHz) δ 8.61 (d, J = 5.2 Hz, 1H), 8.32 (s, 1H), 8.22 (d, J = 8.2 Hz, 1H), 7.88 (d, J = 7.6 Hz, 1H), 7.66 (t, J = 7.9 Hz, 1H), 7.41 (d, J = 5.2 Hz, 1H), 6.19 (s, 1H),2.52 (s, 3H).

INTERMEDIATE COMPOUNDS 108 and 109:

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INTERMEDIATE COMPOUND 108

INTERMEDIATE COMPOUND 109

An ethanol (150mL) solution of INTERMEDIATE COMPOUND

15 107 (5.0g, 13mmol) and 2-aminopyrimidine (5.1g, 53mmol) was heated at reflux for 18h under argon. The contents of the reaction flask were cooled and concentrated *in vacuo*. Water and sat. NaHCO₃ (aq.) were added and the resulting mixture was

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extracted with methylene chloride (3×). The combined organic extracts were dried with Na₂SO₄ (anh.), filtered, and concentrated *in vacuo*. The crude product was subjected to flash column chromatography (methylene chloride methanol 99:1). Two products co-eluted. Product containing fractions were combined and the solvent removed in vacuo to give a tan solid which was triturated with ether and filtered to give a white solid, **INTERMEDIATE COMPOUND 108** (1.24g). The filtrate was concentrated in vacuo and rechromatographed using hexane ethyl acetate 30:70 to give after evaporation a tan foam, **INTERMEDIATE COMPOUND 109** (2.32g).

INTERMEDIATE COMPOUND 108: ¹H NMR (CDCl₃, 300MHz)

8 8.68 (m, 1H), 8.58 (d, 1H), 8.15 (m, 1H), 8.02 (d, 1H), 7.80 (s, br, 2H), 7.70 (m, 2H), 6.92 (m, 1H), 1.84 (s, 3H).

INTERMEDIATE COMPOUND 109: ¹H NMR (CDCl₃, 300MHz) δ 9.88 (dd, J = 6.9, 2.1 Hz, 1H), 8.74 (dd, J = 4.1, 2.1 Hz, 1H), 8.36 (d, J = 5.4 Hz, 1H), 8.06 (s, 1H), 7.91 (d, J = 7.8 Hz, 1H), 7.72 (d, J = 7.8 Hz, 1H), 7.59 (t, J = 7.7 Hz, 1H), 7.09 (dd, J = 6.9, 3.9 Hz, 1H), 6.88 (d, J = 5.4 Hz, 1H), 2.65 (s, 3H).

INTERMEDIATE COMPOUND 110:

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INTERMEDIATE COMPOUND 110 was prepared using

INTERMEDIATE COMPOUND 108 by a procedure like that described for the preparation of INTERMEDIATE COMPOUND 10 above in Scheme 2.

 1 H NMR (CDCl₃, 300MHz) δ 9.00 (d, 1H), 8.75 (m, 1H), 8.56 (d, 1H), 8.18 (m, 1H), 2.73 (s, 3H).

25 **EXAMPLE 126 (COMPOUND 111):**

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EXAMPLE 126 was prepared using INTERMEDIATE

COMPOUND 110 by a procedure like that described above for the preparation of

COMPOUND 11 (EXAMPLE A01).

MS (M+H) m/z 461

INTERMEDIATE COMPOUND 112:

INTERMEDIATE COMPOUND 112 was prepared using

INTERMEDIATE COMPOUND 109 by a procedure like that described for the preparation of INTERMEDIATE COMPOUND 10 above in Scheme 2.

MS (M+H) m/z 420

EXAMPLES 127-130 in TABLE 5 below were prepared by reacting

15 INTERMEDIATE COMPOUND 112 with an amine using a procedure like that described above for the preparation of EXAMPLE A01 (COMPOUND 11).

TABLE 5

EXAMPLE	R	MS (M+H) m/z
127	s.₹ Ph	461
128	see 🔷	425
129	z.e.	361
130	Σ ² CI	495

5 **Scheme 11:**

INTERMEDIATE COMPOUND 117:

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INTERMEDIATE COMPOUND 117 was prepared from INTERMEDIATE COMPOUND 107 and 2-aminopyridine using a procedure like that described for the preparation of INTERMEDIATE COMPOUND 9.

 $^1 H$ NMR (CDCl₃, 300MHz) δ 9.55 (m, 1H), 8.32 (d, 1H), 8.00 (s, 1H), 7.00 (m, 1H), 6.80 (d, 1H), 2.62 (s, 3H).

INTERMEDIATE COMPOUND 118:

10 INTERMEDIATE COMPOUND 118 was prepared using INTERMEDIATE COMPOUND 117 by a procedure like that described for the preparation of INTERMEDIATE COMPOUND 10.

 1H NMR (CDCl₃, 300MHz) δ 9.90 (d, 1H), 8.55 (d, 1H), 7.99 (s, 1H), 7.28 (m, 1H), 7.15 (m, 1H), 3.42 (s, 3H).

EXAMPLES 131-134 in **TABLE 6** were prepared by reacting **INTERMEDIATE COMPOUND 118** with an amine using a procedure like that described for the preparation of **EXAMPLE A01** (**COMPOUND 11**).

TABLE 6

		DLE V	
EXAMPLE	R	¹ H NMR (CDCl ₃ ,	MS (M+H)
	<u> </u>	300MHz) δ	nv/z
131	Ph		460
132	15 O	8.14 (m, 1H), 8.03 (s, br, 1H), 7.75-7.65 (m, 2H), 7.55 (m, 1H), 7.36 (m, 1H),	
133	.5~	9.50 (d, 1H), 8.15 (d, 1H), 8.04 (s, 1H), 7.85 (d, 1H), 6.94 (m, 1H), 6.40 (d, 1H),	
134	H N N N N N N N N N N N N N N N N N N N	9.45 (d, 1H), 8.15 (d, 1H), 8.03 (s, 1H), 7.85 (d, 1H), 7.75-7.65 (m, 2H), 7.53 (m, 1H), 7.39 (m, 1H), 6.95 (m,	

EXAMPLE 135 (COMPOUND 123):

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An ethyl acetate (100mL) solution of **EXAMPLE 134** (0.50g, 1.02mmol) was cooled in an ice bath with stirring. Hydrogen chloride gas was bubbled through the solution for 5min. After 15min. the solvent was removed in vacuo and the remaining solid was recrystallized from acetonitrile to give **EXAMPLE 135** as a solid (84mg).

 1 H NMR (DMSO-d₆, 300MHz) δ 8.33 (d, 1H), 8.09-7.95 (m, 5H), 7.83 (m, 1H), 7.54 (m, 1H), 3.70 (m, 2H), 3.11 (m, 2H).

Scheme 12:

EXAMPLES 136-138 in **TABLE 7** were prepared using a synthetic sequence like that described for the preparation of compounds in **TABLE 6** except 2-amino-4-picoline was used in the place of 2-aminopyridine in the initial condensation reaction with compound **INTERMEDIATE COMPOUND 107**.

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TABLE 7

EXAMPLE	R	¹ H NMR (CDCl ₃ , 300MHz) δ	MS (M+H) m/z	
136	y.s. Ph	8.10 (d, 1H), 7.95 (s, 1H), 7.80 (d, 1H), 7.64 (m, 1H), 7.53-7.30 (m, 8H), 6.38 (m,		
137	3. ² \\ \\ \\ \\ \\ \\ \\ \\ \\ \\ \\ \\ \\	9.40 (m, 1H), 8.13 (d, 1H), 8.03 (s, 1H), 7.85 (d, 1H), 7.65 (d, 1H), 7.52 (m, 2H), 6.79 (m, 1H), 6.40 (d, 1H),		

EXAMPLE	R	¹ H NMR (CDCl ₃ , 300MHz) δ	MS (M+H) m/z
138	N O	9.37 (d, 1H), 8.10 (d, 1H), 8.00 (s, 1H), 7.83 (m, 1H),	
		7.65 (m, 1H), 7.50 (m, 2H), 6.79 (m, 1H), 6.40 (d, 1H),	

Scheme 13:

EXAMPLE 139:

5 **EXAMPLE 139** was prepared using a synthetic sequence like that described for the preparation of the compounds in **TABLE 6** except 2-amino-5-picoline was used in the place of 2-aminopyridine in the initial condensation reaction with **INTERMEDIATE COMPOUND 107**.

¹H NMR (CDCl₃, 300MHz) δ 8.95 (s, br, 1H), 8.11 (d, 1H), 8.00 (s, 1H), 7.81 (d, 1H), 7.65-7.55 (m, 2H), 7.52-7.42 (m, 3H), 7.38 (m, 1H), 7.29 (m, 1H), 7.15 (m, 1H), 6.41 (d, 1H), 5.81 (m, 1H), 5.30 (m, 1H), 2.20 (s, br, 3H), 1.61 (d, 3H).

Scheme 14:

EXAMPLE 140:

EXAMPLE 140 was prepared using a synthetic sequence like that described for the preparation of compounds in **TABLE 6** except 2-amino-3-picoline was used in the place of 2-aminopyridine in the initial condensation reaction with compound **INTERMEDIATE COMPOUND 107**.

MS (M+H) m/z 474

Scheme 15:

INTERMEDIATE COMPOUND 129:

HO NH₂

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INTERMEDIATE COMPOUND 129 was prepared using a synthetic sequence similar to that described for the preparation of INTERMEDIATE COMPOUND 5 except 6-aminonicotinic acid was used in the place of INTERMEDIATE COMPOUND 3.

¹H NMR (CD₃OD, 300MHz) δ 7.85 (m, 1H), 7.48 (m, 1H), 6.59 (m, 1H), 4.42 (s, 2H).

EXAMPLE 141 (COMPOUND 130):

EXAMPLE 141 was prepared using a synthetic sequence like that described in **Scheme 10** for the preparation of **EXAMPLE 127** except

5 INTERMEDIATE COMPOUND 129 was used in the place of 2-aminopyrimidine in the condensation reaction with compound INTERMEDIATE COMPOUND 107.

¹H NMR (CDCl₃, 300MHz) δ 9.10 (m, 1H), 8.15 (d, 1H), 8.00 (s, 1H), 7.82 (d, 1H), 7.65 (m, 1H), 7.54-7.30 (m, 7H), 6.42 (d, 1H), 5.60 (m, 1H), 5.25 (m, 1H), 4.45 (s, br, 1H), 1.62 (d, 3H).

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EXAMPLE 142 (COMPOUND 131):

Molecular sieves (4Å) were added to a methylene chloride (3mL) solution of **EXAMPLE 141** (50mg, 0.10mmol) under argon. After 5min 4-methylmorpholine *N*-oxide (18mg, 0.15mmol) and tetrapropylammonium

perruthenate (4mg, 0.10mmol) were added. After 1h the contents of the reaction flask were subjected to flash column chromatography purification (hexane ethyl acetate 1:1) to give **EXAMPLE 142** as a white solid (28mg).

MS (M+H) m/z 488

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EXAMPLE 143 (COMPOUND 132):

EXAMPLE 143 was prepared by reductive amination of **EXAMPLE** 142 with benzylamine using a procedure like that described for the preparation of compound 14 (**EXAMPLE A04**).

MS (M+H) m/z 579

Scheme 16:

EXAMPLE 144 (COMPOUND 133):

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EXAMPLE 144 was prepared using a synthetic sequence like that described in **Scheme 10** for the preparation of **EXAMPLE 127** except methyl 6-aminonicotinate was used in the place of 2-aminopyrimidine in the condensation reaction with compound **INTERMEDIATE COMPOUND 107**.

MS (M+H) m/z 518

EXAMPLE 145 (COMPOUND 134):

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A solution of lithium hydroxide (7mg, 0.29mmol) in a minimum amount of water was added to a THF (1.5mL) solution of **EXAMPLE 144** (140mg, 0.27mmol) under argon. After 1.5h THF was removed in vacuo and the contents of the reaction flask were acidified with 1N hydrochloric acid. A solid appeared which was isolated by vacuum filtration. Toluene was added to the solid, stirred, then

removed in vacuo. The remaining solid was triturated with ether and isolated to give **EXAMPLE 145** (60mg).

MS (M+H) m/z 504

5 EXAMPLE 146 (COMPOUND 135):

Triethylamine (0.122mL, 0.88mmol) was added with stirring to a N,N-dimethylformamide (5mL) solution of EXAMPLE 145 (400mg, 0.794mmol), 1-[3-(dimethyamino)propyl]-3-ethylcarbodiimide hydrochloride (167mg, 0.874mmol), 1-hydroxy-7-azabenzotriazole (118mg, 0.874mmol), and piperidine (0.087mL, 0.874mmol) under argon. After 48h the contents of the reaction flask were poured into water and the resulting mixture was extracted with ethyl acetate (3x). The combined organic extracts were dried with Na₂SO₄ (anh.), filtered, and concentrated in vacuo. The crude product was subjected to flash column chromatography (ethyl acetate) to give after evaporation a yellow oil, (360mg).

MS (M+H) m/z 571

EXAMPLE 147 (COMPOUND 136):

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EXAMPLE 147 was prepared from **EXAMPLE 145** using a procedure like that described for compound **EXAMPLE 146** except replacing piperidine with *N,N*-dimethylenediamine.

MS (M+H) m/z 574

EXAMPLE 148 (COMPOUND 137):

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Diphenylphophoryl azide (0.130mL, 0.596mmol) and benzyl alcohol (0.093mL, 0.891mmol) were added to a toluene (5mL) solution of 134 (150mg, 0.297mmol) and the resulting solution was heated at reflux 24h. The contents of the reaction flask were cooled and the solvent removed in vacuo. The remaining residue was subjected to flash column chromatography (ethyl acetate hexane 30:70 then 50:50) to give after evaporation **EXAMPLE 148** (96mg).

MS (M+H) m/z 609

EXAMPLE 149 (COMPOUND 138):

EXAMPLE 149 was prepared from **EXAMPLE 141** using a procedure like that described for the preparation of **EXAMPLE A02** (**COMPOUND 12**).

MS (M+H) m/z 515

EXAMPLE 150 (COMPOUND 139):

10 EXAMPLE 150 was prepared from EXAMPLE 149 using a procedure like that described for the preparation of EXAMPLE A03 (COMPOUND 13).

MS (M+H) m/z 489

15 **EXAMPLE 151 (COMPOUND 140):**

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EXAMPLE 151 was prepared from **EXAMPLE 150** by reductive amination with formaldehyde using a procedure like that described for the preparation of **EXAMPLE A04 (COMPOUND 14)**.

MS (M+H) m/z 517

EXAMPLE 152 (COMPOUND 141):

A methylene chloride (3mL) solution of **EXAMPLE 150** (25mg, 0.05mmol) was cooled in an ice bath under argon. Methanesulfonyl chloride (0.020mL, 0.263mmol) and triethylamine (0.041mL, 0.297mmol) were added and the reaction was allowed to warm to room temperature. The solvent was removed in vacuo and the remaining residue was subjected to flash column chromatography (ethyl acetate) to afford after evaporation a white solid 141 (15mg).

MS (M+H) m/z 567

Scheme 17:

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INTERMEDIATE COMPOUND 142

A methanol (250mL) solution of 2-aminonicotinic acid (10.5g, 76mmol), and sulfuric acid (20mL) was refluxed 18h. The reaction was cooled to room temperature and the solvent was removed in vacuo. Sat. sodium bicarbonate (aq.) was added and the mixture was extracted with ethyl acetate. The organic layer was dried with anhydrous sodium sulfate, filtered, and evaporated in vacuo to give a white solid INTERMEDIATE COMPOUND 142 (5.5g).

 1 H NMR (CD₃OD, 300MHz) δ 8.19 (m, 2H), 7.64 (m, 1H), 3.89 (s, 3H).

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INTERMEDIATE COMPOUND 143:

INTERMEDIATE COMPOUND 143 was prepared using a synthetic sequence like that described in Scheme 10 for the preparation of INTERMEDIATE COMPOUND 112 except INTERMEDIATE COMPOUND 142 was used in the place of 2-aminopyrimidine in the condensation reaction with compound INTERMEDIATE COMPOUND 107.

¹H NMR (CDCl₃, 300MHz) δ 10.10 (m, 1H), 8.58 (d, 1H), 8.25 (m, 1H), 8.00 (s, 1H), 7.85 (m, 1H), 7.76 (m, 1H), 7.63 (m, 1H), 7.26 (m, 2H), 4.09 (s, 3H), 3.43 (s, 3H).

EXAMPLES 153 (COMPOUND 144) and 154 (COMPOUND 145):

EXAMPLE 153 (COMPOUND 144)

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EXAMPLE 154 (COMPOUND 145)

S-(-)- α -Methylbenzylamine (15mL) and INTERMEDIATE

- 10 COMPOUND 143 (2.61g, 5.48mmol) were combined under argon and heated at 60°C for 1h. Cooled, added citric acid (aq.) and extracted with ethyl acetate. Dried the organic layer with anhydrous sodium sulfate and removed solvent in vacuo to give a yellow solid. Flash column chromatography (ethyl acetate hexane 25:75) followed by reverse phase preparative HPLC afforded, after evaporation, **EXAMPLES 153** and
- 15 **EXAMPLES 154**.

EXAMPLE 153: MS (M+H) m/z 518 **EXAMPLE 154**: MS (M+H) m/z 607

EXAMPLE 155 (COMPOUND 146):

Lithium aluminum hydride solution (1mL, 1M) was slowly added to a THF solution of EXAMPLE 154 (300mg, 0.580mmol) under argon at room temperature. After 18h the reaction was quenched with water and sodium hydroxide (aq.). Magnesium sulfate was added and the mixture was filtered. The filtrate was evaporated in vacuo to give a red oil. Flash column chromatography gave

EXAMPLE 155 after evaporation as a white solid (60mg).

MS (M+H) m/z 490

EXAMPLE 156 (COMPOUND 147):

EXAMPLE 156 was prepared from EXAMPLE 155 using a procedure like that described for the preparation of EXAMPLE A02 (COMPOUND 12).

EXAMPLE 157 (COMPOUND 148):

5 EXAMPLE 157 was prepared from EXAMPLE 156 using a procedure like that described for the preparation of EXAMPLE A03 (COMPOUND 13).

Combustion analysis for **EXAMPLE 157**: Calculated for C₂₇H₂₃N₆F₃.0.05H₂0.0.45MeOH C 65.43%, H 4.98%, N 16.68% Found: C 65.43%, H 4.62%, N 16.61%.

EXAMPLE 158 (COMPOUND 149):

EXAMPLE 158 was prepared from EXAMPLE 157 using a procedure like that described for the preparation of EXAMPLE 152.

MS (M+H) m/z 567

EXAMPLE 159 (COMPOUND 150):

EXAMPLE 159 was prepared from EXAMPLE 153 using a procedure like that described for the preparation of compound EXAMPLE 145.

Combustion analysis for EXAMPLE 159: Calculated for C₂₇H₂₀N₅O₂F₃.0.10H₂0.1.95TFA C 51.00%, H 3.07%, N 9.63% Found: C 51.00%, H 3.04%, N 9.62%.

EXAMPLE 160 (COMPOUND 151):

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A neat solution of **COMPOUND 144** in *N,N*-dimethylethylenediamine was heated at 80°C under argon for 2h. The contents of the reaction flask were cooled to room temperature and acetonitrile/water/methanol was added. The resulting solution was subjected to reverse phase preparative HPLC to give after evaporation **COMPOUND 151**.

MS (M+H) m/z 574

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EXAMPLE 161 (COMPOUND 152):

A neat solution of 4-amino-1-BOC-piperidine (300mg, 1.5mmol) and 144 (50mg, 0.1mmol) was heated at 80°C under argon for 18h. cooled to room temperature and added ethyl acetate and water. The organic layer was dried with anhydrous sodium sulfate, filtered and concentrated in vacuo. Methylene chloride (4mL) and trifluoroacetic acid (4mL) were added. After 3h the solvents were evaporated in vacuo and the remaining residue was subjected to reverse phase preparative HPLC to give after evaporation a yellow solid, 152 (10mg, 17%).

MS (M+H) m/z 586

Scheme 18:

INTERMEDIATE COMPOUND 153:

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INTERMEDIATE COMPOUND 153 was prepared using a synthetic sequence like that described in Scheme 10 for the preparation of INTERMEDIATE

COMPOUND 112 except INTERMEDIATE COMPOUND 5 was used in the place of 2-aminopyrimidine in the condensation reaction with compound INTERMEDIATE COMPOUND 107.

INTERMEDIATE COMPOUND 153: ¹H NMR (CDCl₃, 300MHz) 5 δ 9.81 (d, 1H), 8.55 (d, 1H), 7.95 (s, 1H), 7.80 (m, 3H), 7.65 (m, 1H), 7.25 (m, 1h), 7.11 (m, 1H), 4.86 (s, 2H), 3.41 (m, 2H).

EXAMPLE 162 (COMPOUND 154):

10 EXAMPLE 162 was prepared from INTERMEDIATE

COMPOUND 153 using a procedure like that described for the preparation of EXAMPLE A01 (COMPOUND 11) except 3-ethoxypropylamine was used in the place of s-(-)-\alpha-methylbenzylamine.

MS (M+H) m/z 472

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EXAMPLE 163 (COMPOUND 155):

EXAMPLE 163 was prepared from INTERMEDIATE

COMPOUND 153 using a procedure like that described for the preparation of EXAMPLE A01 (COMPOUND 11) except N,N-ethylenediamine was used in the place of s-(-)-α-methylbenzylamine.

MS (M+H) m/z 457

EXAMPLE 164 (COMPOUND 156) and INTERMEDIATE COMPOUND 157:

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EXAMPLE 164 (COMPOUND 156)

INTERMEDIATE COMPOUND 157

S-(-)- α -Methylbenzylamine (5mL) was added to an isopropanol

(15mL) solution of INTERMEDIATE COMPOUND 153 (1.90g, 4.24mol) under argon and the resulting mixture was heated at 60°C 18h. The contents of the reaction flask were cooled to room temperature and treated with citric acid (aq.). The pH was adjusted to 4.5 with NaOH (aq.) and extracted with ethyl acetate (2×). The combined organic extracts were dried with anhydrous sodium sulfate, filtered, and the filtrate concentrated in vacuo to give a red oil. Flash column chromatography (ethyl acetate hexane 40:60 then 70:30) to give two portions after evaporation: 1. 156 (349mg) and 2. A mixture (610mg) of 156 and 157.

156: 1 H NMR (CDCl₃, 300MHz) δ 8.11 (d, 1H), 7.95 (s, 1H), 7.80 (d, 1H), 7.70-7.30 (m, 10H), 6.36 (d, 1H), 5.68 (m, 1H), 5.16 (m, 1H), 4.72 (m, 2H), 1.63 (d, 3H).

5 EXAMPLE 166 (COMPOUND 158) and INTERMEDIATE COMPOUND 159:

EXAMPLE 166 (COMPOUND 158)

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INTERMEDIATE COMPOUND 159

Diphenylphophoryl azide (0.0.323mL, 1.50mmol) and 1,8-diazabicyclo[4.5.0]undec-7-ene (0.224mL, 1.50mmol) were added to a toluene (5mL) solution of **EXAMPLE 141** (115mg, 0.235mmol) under argon. After 18h the reaction was poured into water and extracted (3×) with ethyl acetate. The combined organic portions were dried with Na₂SO₄ (anh.), filtered, and concentrated in vacuo. The crude product was subjected to flash column chromatography (ethyl acetate hexane 10:90 then 20:80) to give after evaporation two products: 1. White solid, **158** (100mg) and 2. Solid, **159** (38mg).

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158: 1 H NMR (CDCl₃, 300MHz) δ 8.15 (d, 1H), 7.97 (s, 1H), 7.83 (d, 1H), 7.66 (d, 1H), 7.59-7.33 (m, 9H), 5.70 (m, 1H), 5.16 (m, 1H), 4.40 (s, 2H), 1.65 (d, 3H).

159: ¹H NMR (CDCl₃, 300MHz) δ 9.60 (d, 1H), 8.32 (d, 1H), 8.00 (s, 1H), 7.82 (d, 1H), 7.70 (m, 2H), 7.57 (m, 1H), 6.95 (m, 1H), 6.75 (m, 1H), 5.39 (m, 1H), 4.50 (s, 2H), 1.50 (d, 6H).

INTERMEDIATE COMPOUND 160:

10 INTERMEDIATE COMPOUND 160 was prepared from INTERMEDIATE COMPOUND 159 using a procedure like that described for the preparation of EXAMPLE A03 (COMPOUND 13).

MS (M+H) m/z 428

15 **EXAMPLE 169 (COMPOUND 161):**

EXAMPLE 169 was prepared from EXAMPLE 166 using a procedure like that described for the preparation of EXAMPLE A03 (COMPOUND 13).

MS (M+H) m/z 489

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EXAMPLE 170 (COMPOUND 162):

EXAMPLE 170 was prepared from EXAMPLE 169 using a procedure like that described for the preparation of EXAMPLE 152 (COMPOUND 10 141).

MS (M+H) m/z 567

EXAMPLE 171 (COMPOUND 163):

EXAMPLE 171 was prepared from **EXAMPLE 169** by reductive amination using a procedure like that described for the preparation of **EXAMPLE A04 (COMPOUND 14)**.

MS (M+H) m/z 517

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EXAMPLE 172 (COMPOUND 164):

EXAMPLE 172 was prepared from EXAMPLE 169 by reductive amination using a procedure like that described for the preparation of EXAMPLE A04 (COMPOUND 14) except formaldehyde (aq.) was replaced with acetone.

MS (M+H) m/z 531

EXAMPLE 173 (COMPOUND 165):

15 **EXAMPLE 173** was prepared from **EXAMPLE 169** by reductive amination using a procedure like that described for the preparation of **EXAMPLE**A04 except formaldehyde (aq.) was replaced with ethyl formate.

MS (M+H) m/z 517

EXAMPLE 174 (COMPOUND 166):

A THF solution of **EXAMPLE 173** (61mg, 0.12mmol) under argon was treated with 1M borane-THF solution (0.59mL, 0.59mmol) and stirred at room temperature. After 18h, 2M hydrochloric acid was added and after 2h the reaction was made basic with sat. sodium bicarbonate (aq.). The resulting mixture was extracted with ethyl acetate (2×). The combined organic extracts were dried with anhydrous sodium sulfate, filtered, and evaporated in vacuo to give a yellow oil. Flash column chromatography (methylene chloride methanol ammonium hydroxide 95:5:0.5 gave after evaporation a yellow solid of **EXAMPLE 173** (31mg).

MS (M+H) m/z 503

15 Compounds in **TABLE 8** below were prepared by reacting **EXAMPLE 169** with a carboxylic acid using a coupling procedure like that described for the preparation of **EXAMPLE 146**.

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TABLE 8

EXAMPLE	R_	MS (M+H) m/z
175	**\^S\\	653
176	34 A	557
177	F	629
178		697
179	N N	596
180	3-	625

Scheme 19:

5 **EXAMPLE 181:**

EXAMPLE 181 was prepared using a synthetic sequence like that described in Scheme 10 for the preparation of EXAMPLE 127 except INTERMEDIATE COMPOUND 4 was used in the place of 2-aminopyrimidine in the condensation reaction with compound INTERMEDIATE COMPOUND 107.

MS (M+H) m/z 532

EXAMPLE 182 (COMPOUND 174):

15 **EXAMPLE 182** was prepared from **EXAMPLE 181** using a procedure like that described for the preparation of **EXAMPLE 145**.

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 $^{1}\text{H NMR (CD}_{3}\text{OD, 300MHz})~\delta~8.18$ (s, 1H), 8.10 (d, 1H), 7.90 (s, 1H), 7.85 (m, 1H), 7.75 (m, 1H), 7.65 (m, 1H), 7.48-7.25 (m, 7H), 6.30 (d, 2H), 1.60 (d, 3H).

5 Compounds in **TABLE 9** below were prepared by reacting **EXAMPLE 182** with an amine using a coupling procedure like that described for the preparation of **EXAMPLE 146**.

10

	TABLE 9	
EXAMPLE	R	MS (M+H) m/z
183	N.	571
184	H _{2g} N O	589
185	34 N_	531
186	THE PART OF THE PA	472

EXAMPLE	R	MS (M+H) m/z
187	H F	625

Other **EXAMPLES** of the invention are shown in the following **TABLE 10**. These **EXAMPLES** are made similarly to the compounds and **Schemes** shown above.

. 5

TABLE 10

	CADLE IV	
Example	M+1	H-NMR: (400MHz)δ
B001	467.4	CDCl ₃ : 8.60(br, 1H), 8.38(s,
F Chiral		1H), 8.10(d, J=5.5Hz, 1H),
		7.63(m, 2H), 7.45(m, 6H),
		7.12(m, 2H), 6.53(br, 1H),
		6.42(d, J=5.3Hz, 1H),
H H		6.12(br, 1H), 5.84(br, 1H),
N N N N N N N N N N N N N N N N N N N		5.16(m, 1H), 4.56(d,
HN O		J=6.3Hz, 2H), 1.63(d,
	1	J=7.0Hz, 3H)

Example	M+1	H-NMR: (400MHz)δ
B002 Chiral N N N OH OH	422.3	CD ₃ OD: 8.58(br, 1H), 8.04(d, J=5.3Hz, 1H), 7.56(m, 3H), 7.42(m, 4H), 7.28(m, 1H), 7.18(m, 2H), 6.82(br, 1H), 6.27(d, J=5.3Hz, 1H), 5.11(m, 1H), 4.01(s, 2H),3.68(m, 4H), 2.82(m, 1H), 1.56(d, J=7.1Hz, 3H)
B003 Chiral	453.3	CDCl3: 8.56(br, 1H), 8.11(d, J=5.3Hz, 1H), 7.42(m, 7H), 7.24(m, 1H), 7.18(m, 2H), 6.66(br, 1H), 6.22(d, J=5.3Hz, 1H), 5.10(m, 1H), 3.78(s, 2H), 2.41(s, 3H), 1.58(d, J=7.0Hz, 3H)

Brown le	136.1	
Example	M+1	H-NMR: (400MHz)δ
B004 F. Chiral	481.3	CD3OD: 8.56(br, 1H),
Criirai		8.02(d, J=5.3Hz, 1H),
		7.56(m, 3H), 7.42(m, 5H),
		7.28(m, 1H), 7.14(m, 2H),
		6.76(br, 1H), 6.28(d,
		J=5.3Hz, 1H), 5.12(m, 1H),
		3.85(s, 2H), 2.88(m, 1H),
HN		1.57(d, J=7.0Hz, 3H),
		1.25(m, 6H)
		-
B005	497.4	CDCl3: 8.62(br, 1H), 8.12(d,
Chiral		J=5.3Hz, 1H), 7.64(m, 2H),
		7.45(m, 6H), 7.13(m, 2H),
		6.62(br, 1H), 6.42(d,
		J=5.3Hz, 1H), 5.62(m, 1H),
		5.18(m, 1H), 3.71(t, J=5.2Hz,
OH OH		2H), 3.67(s, 2H), 2.70(t,
HN		J=5.3Hz, 2H), 2.34(s, 3H),
		1.63(d, J=7.1Hz, 3H)
B006	537.4	CD2OD, 9.564- IID
F, Chiral	JJ 7.4	CD3OD: 8.56(br, 1H), 8.04(d, J=5.3Hz, 1H),
		7.57(m, 3H), 7.42(m, 4H),
		I
<i>}_\(\)</i>		7.26(m, 1H), 7.18(m, 2H),
		6.82(br, 1H), 6.28(d,
N N		J=5.3Hz, 1H), 5.12(m, 1H),
HN		4.16(m, 1H), 3.80(m, 4H),
		2.62(m, 2H), 2.40(s, 3H),
		2.02(m, 4H), 1.56(d,
~		J=7.1Hz, 3H)
	· · · · · · · · · · · · · · · · · · ·	

T 1.	26.1	TT AD ED (4007 FT) S
Example	M+1	H-NMR: (400MHz)δ
B007	538.5	CD3OD: 8.42(br, 1H),
\		8.05(d, J=5.2Hz, 1H),
· · · · · · · · · · · · · · · · · · ·		7.52(m, 2H), 7.40(m, 5H),
N		7.24(m, 1H), 6.62(br, 1H),
N	la control de la	6.29(d, J=5.3Hz, 1H),
N N N		5.10(m, 1H), 4.62(s, 2H),
		2.97(t, J=6.6H 2H), 2.62(t,
		J=6.5z, 2H), 2.15(s, 6H),
		1.56(d, J=7.1Hz, 3H)
B008	510.4	CD3OD: 8.42(br, 1H),
F		8.01(d, J=5.2Hz, 1H),
		7.52(m, 2H), 7.40(m, 5H),
	1	7.24(m, 1H), 6.62(br, 1H),
		6.21(d, J=5.3Hz, 1H),
N N N		5.07(m, 1H), 4.45(s, 2H),
N N N N N N N N N N N N N N N N N N N	Í	3.10(t, J=6.7Hz, 2H), 2.61(t,
		J=6.6Hz, 2H), 1.54(d,
		J=6.8Hz, 3H)
	495.4	CDCl3: 8.72(br, 1H), 8.10(d,
B009		J=5.3Hz, 1H), 7.61(m, 3H),
	-	7.46(m, 4H), 7.36(m, 1H),
		7.11(m, 2H), 6.72(br, 1H),
		6.42(d, J=5.3Hz, 1H),
N		5.61(br, 1H), 5.20(m, 1H),
N !		3.60(br, 2H), 2.98(br, 1H),
N N		2.23(br, 3H), 1.64(d,
\ \ \		J=7.0Hz, 3H), 1.27(br, 6H)
		-

Example	M+1	H-NMR: (400MHz)δ
B010	538.5	CD3OD: 8.54(br, 1H),
Ę		8.03(d, J=5.1Hz, 1H),
		7.55(m, 2H), 7.48(s, 1H),
		7.42(m, 4H), 7.24(m, 1H),
> −N		7.18(m, 2H), 6.78(br, 1H),
		6.25(d, J=5.2Hz, 1H),
		5.12(m, 1H), 3.57(s, 2H),
N N		2.42(m, 4H), 2.27(s, 6H),
	·	1.77(m, 2H), 1.56(d,
		J=7.1Hz, 3H)
		3,
B011	574.4	CD3OD: 8.56(br, 1H),
F		8.04(d, J=5.3Hz, 1H),
		7.56(m, 2H), 7.52(s, 1H),
		7.43(m, 4H), 7.28(m, 1H),
		7.18(m, 2H), 6.81(br, 1H),
N N N N O O	1	6.30(d, J=5.2Hz, 1H),
N N N N N N N N N N N N N N N N N N N		5.12(m, 1H), 3.65(s, 2H),
		3.27(t, J=6.4Hz, 2H), 2.97(s,
		3H), 2.64(t, J=6.4Hz, 2H),
		2.31(s, 3H), 1.56(d, J=7.0Hz,
	·	3H)
B012	536.4	CD3OD: 8.58(br, 1H),
		8.04(d, J=5.3Hz, 1H),
		7.56(m, 2H), 7.52(s, 1H),
		7.43(m, 4H), 7.28(m, 1H),
		7.18(m, 2H), 6.78(br, 1H),
N N I		6.28(d, J=5.3Hz, 1H),
N N N		5.12(m, 1H), 3.60(s, 2H),
		3.40(m, 4H), 2.76(t, J=6.7Hz,
		2H), 2.47(t, J=6.7Hz, 2H),
		2.27(s, 3H), 2.17(m, 2H),
		1.57(d, J=7.1Hz, 3H)

	r	<u> </u>
Example	M+1	H-NMR: (400MHz)δ
B013	527.4	CDCl3: 8.58(br, 1H), 8.10(d,
T		J=5.3Hz, 1H), 7.64(m, 2H),
		7.45(m, 6H), 7.13(m, 2H),
		6.60(br, 1H), 6.41(d,
N		J=5.3Hz, 1H), 5.80(br, 1H),
N N		5.12(m, 1H), 3.90(s, 2H),
N N		3.74(m, 4H), 3.14(m, 1H),
		2.38(s, 3H), 2.20(br, 2H),
0		1.62(d, J=7.0Hz, 3H)
B014	433.4	CD3OD: 9.62(br, 1H),
N N		8.05(d, J=5.4Hz, 1H),
N N N		7.61(m, 3H), 7.21(m, 3H),
		6.31(d, J=5.3Hz, 1H), 3.74(s,
, N		2H), 3.38(s, 2H), 2.41(s, 6H),
()		1.01(s, 9H)
)==/ F	•	
B015	447.3	CD3OD: 9.61(br, 1H),
		8.05(d, J=5.4Hz, 1H),
7 N		7.61(m, 3H), 7.21(m, 3H),
N—		6.31(d, J=5.3Hz, 1H), 3.73(s,
N N	·	2H), 3.36(s, 2H), 2.61(m,
> N<	-	2H), 2.34(s, 3H), 1.18(t,
NIN		J=7.3Hz, 3H), 1.01(s, 9H)
F		

	<u> </u>	
Example	M+1	H-NMR: (400MHz)δ
B016	552.3	CD3OD: 8.57(br, 1H),
1		8.06(d, J=5.3Hz, 1H),
		7.57(m, 2H), 7.50(s, 1H),
N		7.42(m, 4H), 7.28(m, 1H),
N N		7.18(m, 2H), 6.71(br, 1H),
		6.29(d, J=5.5Hz, 1H),
F-\ \ \ \ \ \ \ \ \ \ \ \ \ \ \ \ \ \ \		3.91(m, 1H), 3.69(m, 2H),
N		3.25(s, 3H), 2.99(s, 3H),
_N		2.26(s, 3H), 1.58(d, J=7.0Hz,
	,	3H), 1.25(d, J=6.6Hz, 3H)
O N		
B017	564.1	CD3OD: 8.58(br, 1H),
, rui]	8.05(d, J≈5.3Hz, 1H),
, N		7.58(m, 2H), 7.51(s, 1H),
		7.42(m, 4H), 7.27(m, 1H),
N		7.18(m, 2H), 6.81(br, 1H),
- N		6.29(d, J=5.3Hz, 1H),
N		5.13(m, 1H), 3.71(s, 2H),
, N		3.55(t, J=6.7Hz, 2H), 3.42(t,
	:	J=6.9Hz, 2H), 3.34(s, 2H),
0 N		2.36(s, 3H), 1.99(m, 2H),
		1.87(m, 2H), 1.57(d,
		J=7.0Hz, 3H)
B018	392.2	CDCl3: 9.31(d, 1H), 8.11(d,
X		1H), 7.64(m, 3H), 7.11(m,
/ N		2H), 6.92(m, 1H), 6.41(d,
$N \longrightarrow N$		1H), 5.26(br, 1H), 4.80(s,
N /		2H), 1.54(s, 9H)
N -		
F		
		<u> </u>

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Example	M+1	H-NMR: (400MHz)δ
B019		
В019	550.1	CD3OD: 8.58(br, 1H),
, in		8.04(d, J=5.3Hz, 1H),
N		7.58(m, 2H), 7.51(s, 1H),
		7.43(m, 4H), 7.25(m, 1H),
N		7.18(m, 2H), 6.81(br, 1H),
l N		6.28(d, J=5.2Hz, 1H),
F-\\\\\\\\\\\\\\\\\\\\\\\\\\\\\\\\\\\\		5.13(m, 1H), 4.30(m, 2H),
		4.04(m, 2H), 3.66(s, 2H),
/N		3.13(s, 2H), 2.32(m, 5H),
0 N		1.57(d, J=7.1Hz, 3H)
B020	550.1	CD3OD: 8.57(br, 1H),
, iii		8.04(d, J=5.3Hz, 1H),
		7.57(m, 2H), 7.48(s, 1H),
N-N		7.43(m, 4H), 7.30(m, 1H),
		7.18(m, 2H), 6.70(br, 1H),
	i	6.28(d, J=5.3Hz, 1H),
		5.12(m, 1H), 3.71(s, 2H),
F		3.65(m, 3H), 2.94(s, 6H),
N		1.57(d, J=7.0Hz, 3H)
, N,	٠.	
.]		
N O		
•		

Example	M+1	H-NMR: (400MHz)δ
Example B021 N N N N N N N N N N N N N N N N N N	M+1 497.1	H-NMR: (400MHz)δ CD3OD: 8.48(br, 1H), 8.02(d, J=5.3Hz, 1H), 7.54(m, 2H), 7.41(m, 5H), 7.23(m, 1H), 7.17(m, 2H), 6.61(br, 1H), 6.23(d, J=5.5Hz, 1H), 5.07(m, 1H), 4.34(s, 2H), 3.72(s, 3H), 1.55(d, J=7.1Hz, 3H)
N O		
F N N N N N N N N N N N N N N N N N N N	580.2	CD3OD: 8.60(br, 1H), 8.05(d, J=5.6Hz, 1H), 7.57(m, 2H), 7.53(s, 1H), 7.43(m, 4H), 7.28(m, 1H), 7.18(m, 2H), 6.80(br, 1H), 6.29(d, J=5.3Hz, 1H), 5.12(m, 1H), 4.14(m, 2H), 3.63(s, 2H), 3.53(m, 4H), 2.49(m, 4H), 1.57(d, J=7.0Hz, 3H), 1.25(m, 3H)
	·	

<u></u>		
Example	M+1	H-NMR: (400MHz)δ
B023	445.2	CD3OD: 9.62(br, 1H),
N	ļ	8.05(d, J=5.3Hz, 1H),
		7.61(m, 2H), 7.59(s, 1H),
N N		7.21(m, 2H), 7.05(m, 1H),
N N		6.31(d, J=5.3Hz, 1H), 3.78(s,
		2H), 3.42(m, 4H), 3.36(s,
N		2H), 2.21(m, 2H), 1.01(s,
		9H)
F		
B024	447.2	CD3OD: 9.62(br, 1H),
N		8.05(d, J=5.2Hz, 1H),
		7.61(m, 3H), 7.21(m, 2H),
N N		7.15(m, 1H), 6.31(d,
N		J=5.3Hz, 1H), 3.97(s, 2H),
		3.00(m, 1H), 1.27(d,
N		J=6.3Hz, 6H), 1.01(s, 9H)
F T		
B025	419.1	CD3OD: 9.42(m, 1H),
_\		8.09(d, J=5.1Hz, 1H),
TN S	• •	7.61(m, 3H), 7.21(m, 2H),
N		7.10(m, 1H), 6.35(d,
N N		J=5.2Hz, 1H), 3.60(s, 2H),
√ /N		2.30(s, 6H), 1.48(s, 9H)
	'	
[·

Example	124.1	Y > D CD (100) CT > D
Example	M+1	H-NMR: (400MHz)δ
B026	412.0	CD3OD: 9.64(d, J=7.3Hz,
	ŀ	1H), 8.21(d, J=5.2Hz, 1H),
N		7.65(m, 5H), 7.32(m, 4H),
_ <u>_</u>		7.05(m, 2H), 6.53(d,
N N O		J=5.3Hz, 1H), 4.74(s, 2H)
N		
N		
F -		·
B027	433.1	CD3OD: 9.42(m, 1H),
	.55.1	8.09(d, 1H), 7.62(m, 3H),
_/ N		
N		7.21(m, 2H), 7.13(m, 1H),
N		6.36(d, 1H), 4.13(s, 2H),
N N N		3.12(m, 1H), 1.48(s, 9H),
N N		1.23(d, 6H)
		·
)	•	
F	Line Control	
B028	405.1	CD3OD: 9.63(d, J=7.0Hz,
N N		1H), 8.04(d, J=5.2Hz, 1H),
		7.62(m, 3H), 7.20(m, 2H),
N N N	.	7.10(m, 1H), 6.30(d,
		J=5.3Hz, 1H), 4.18(m, 1H),
)ñ		3.62(s, 2H), 2.34(s, 6H),
		1.29(d, J=6.4Hz, 6H)
F		

Example	M+1	H-NMR: (400MHz)δ
B029 N N N N N F	424.1	CDCl3: 9.60(br, 1H), 8.08(d, J=5.2Hz, 1H), 7.63(m, 2H), 7.45(m, 6H), 7.11(m, 2H), 6.41(d, J=5.4Hz, 1H), 5.63(m, 1H), 5.21(m, 1H), 2.42(s, 3H), 1.63(d, J=7.1Hz, 3H)
B030	390.1	CDCl3: 9.42(br, 1H), 8.08(d, J=5.1Hz, 1H), 7.66(m, 2H), 7.46(s, 1H), 7.14(m, 2H), 6.78(m, 1H), 6.42(d, J=5.3Hz, 1H), 5.28(br, 1H), 3.38(d, J=6.3Hz, 2H), 2.48(s, 3H), 1.06(s, 9H)
B031	417.1	CD3OD: 9.60(d, J=7.0Hz, 1H), 8.02(d, J=5.3Hz, 1H), 7.61(m, 2H), 7.53(s, 1H), 7.20(m, 2H), 7.02(m, 1H), 6.30(d, J=5.4Hz, 1H), 4.17(m, 1H), 3.74(s, 2H), 3.38(m, 4H), 2.17(m, 2H), 1.28(d, J=6.4Hz, 6H)

Example	M+1	H-NMR: (400MHz)δ
B032 N N N N N N N N N N N N N N N N N N	419.1	H-NMR: (400MHz)8 CD3OD: 9.61(d, J=7.0Hz, 1H), 8.01(d, J=5.3Hz, 1H), 7.60(m, 3H), 7.20(m, 2H), 7.11(m, 1H), 6.28(d, J=5.3Hz, 1H), 4.16(m, 1H), 3.90(s, 2H), 2.90(m, 1H), 1.28(d, J=6.4Hz, 6H), 1.16(d, J=6.2Hz, 6H)
B033	439.1	CD3OD: 9.64(d, J=7.3Hz, 1H), 8.21(d, J=5.2Hz, 1H), 7.65(m, 5H), 7.32(m, 4H), 7.05(m, 2H), 6.53(d, J=5.3Hz, 1H), 4.74(s, 2H)

Other **EXAMPLES** of the invention are shown in the following **TABLE 11**. These **EXAMPLES** are made similarly to the compounds and **Schemes** shown above.

TABLE 11

		
EXAMPLE	M+1	NMR(CDCl3)

EXAMPLE	M+1	NMR(CDCl3)
F C001	479.4	1.6 (d, J=6.9 Hz, 3H), 2.2 (qn, J=7.0 Hz, 2H), 3.3 (t, J=7.0 Hz, 4H), 3.6 (s, 2H), 5.2 (qn J=6.9 Hz, 1H), 5.8 (broad, 1H), 6.4 (d, J=5.3 Hz, 1H), 6.6 (broad, 1H), 7.1 (m, 2H), 7.3 (m, 1H), 7.4 (m, 5H), 7.6 (m, 2H), 8.1 (d, J=5.2 Hz, 1H), 8.7 (broad, 1H).
N N N N N N N N N N N N N N N N N N N	422.3	1.6 (d, J=7.0 Hz, 3H), 1.8 (broad, 4H), 2.6 (broad, 4H), 3.7 (broad, 2H), 5.2 (qn J=7.0 Hz, 1H), 5.6 (broad, 1H), 6.4 (d, J=5.2 Hz, 1H), 6.7 (broad, 1H), 7.1 (m, 2H), 7.3 (m, 1H), 7.4 (m, 5H), 7.6 (m, 2H), 8.1 (d, J=5.2 Hz, 1H), 8.7 (broad, 1H).
N N N N N N N N N	507.4	1.5 (m, 2H), 1.6 (d, J=6.9 Hz, 3H), 1.6-1.8 (broad, 4H), 2.4 (broad, 4H), 3.5 (s, 2H), 5.2 (qn J=6.9 Hz, 1H), 5.6 (broad, 1H), 6.4 (d, J=5.3 Hz, 1H), 6.7 (broad, 1H), 7.1 (m, 2H), 7.3 (m, 1H), 7.4 (m, 4H), 7.6 (m, 3H), 8.1 (d, J=5.3 Hz, 1H), 8.7 (broad, 1H).
N N N N O CO04	509.4	1.6 (d, J=6.9 Hz, 3H), 2.5 (m, 4H), 3.6 (s, 2H), 3.8 (m, 4H), 5.2 (qn J=6.9 Hz, 1H), 5.8 (broad, 1H), 6.4 (d, J=5.2 Hz, 1H), 6.7 (broad, 1H), 7.1 (m, 2H), 7.3 (m, 1H), 7.4 (m, 4H), 7.6 (m, 3H), 8.1 (d, J=5.2 Hz, 1H), 8.7 (broad, 1H).

NMR(CDCi3)	EVANDIE	75.4	
J=5.7 Hz, 4H), 3.4 (s, 6H), 3.5 (t, J=5.8 Hz, 4H), 3.8 (s, 2H), 5.2 (qn J=7.0 Hz, 1H), 5.7 (broad, 1H), 6.4 (d, J=5.3 Hz, 1H), 6.7 (broad, 1H), 7.1 (m, 2H), 7.3 (m, 1H), 7.4 (m, 4H), 7.6 (m, 3H), 8.1 (d, J=5.3 Hz, 1H), 6.7 (broad, 1H). 525.4 1.6 (d, J=6.9 Hz, 3H), 2.4 (s, 3H), 3.4 (s, 2H), 3.8 (m, 5H), 5.2 (qn J=6.8 Hz, 1H), 5.8 (broad, 1H), 6.4 (d, J=5.2 Hz, 1H), 6.7 (broad, 1H), 7.1 (m, 2H), 7.3 (m, 1H), 7.4 (m, 4H), 7.6 (m, 3H), 8.1 (d, J=5.2 Hz, 1H), 8.7 (broad, 1H). C006 497.4 2.2 (s, 3H), 2.7 (2.7, 2H), 3.4 (s, 3H), 3.6 (t, J=5.7 Hz, 2H), 3.7 (s, 2H), 4.8 (d, J=5.9 Hz, 2H), 5.8 (broad, 1H), 6.5 (d, J=5.3 Hz, 1H), 6.8 (broad, 1H), 6.7 (m, 3H), 8.1 (d, 1H), 6.8 (broad, 1H), 7.1 (m, 2H), 7.4 (m, 5H), 7.6 (m, 3H), 8.1 (d, 1H), 6.8 (broad, 1H), 7.1 (m, 2H), 7.4 (m, 5H), 7.6 (m, 3H), 8.1 (d, 1H), 6.8 (broad, 1H), 7.1 (m, 2H), 7.4 (m, 5H), 7.6 (m, 3H), 8.1 (d, 1H), 6.8 (broad, 1H), 7.1 (m, 2H), 7.4 (m, 5H), 7.6 (m, 3H), 8.1 (d, 1H), 6.8 (broad, 1H), 8.1 (d, 1H), 8.2 (d, 1H),	EXAMPLE	M+1	NMR(CDCl3)
J=5.8 Hz, 4H), 3.8 (s, 2H), 5.2 (qn J=7.0 Hz, 1H), 5.7 (broad, 1H), 6.4 (d, J=5.3 Hz, 1H), 6.7 (broad, 1H), 7.1 (m, 2H), 7.3 (m, 1H), 7.4 (m, 4H), 7.6 (m, 3H), 8.1 (d, J=5.3 Hz, 1H), 8.7 (broad, 1H). 525.4 1.6 (d, J=6.9 Hz, 3H), 2.4 (s, 3H), 3.4 (s, 2H), 3.8 (m, 5H), 5.2 (qn J=6.8 Hz, 1H), 5.8 (broad, 1H), 6.4 (d, J=5.2 Hz, 1H), 6.7 (broad, 1H), 7.1 (m, 2H), 7.3 (m, 1H), 7.4 (m, 4H), 7.6 (m, 3H), 8.1 (d, J=5.2 Hz, 1H), 8.7 (broad, 1H). C006 497.4 2.2 (s, 3H), 2.7 (2.7, 2H), 3.4 (s, 3H), 3.6 (t, J=5.7 Hz, 2H), 3.7 (s, 2H), 4.8 (d, J=5.9 Hz, 2H), 5.8 (broad, 1H), 6.5 (d, J=5.3 Hz, 1H), 6.8 (broad, 1H), 7.1 (m, 2H), 7.4 (m, 5H), 7.6 (m, 3H), 8.1 (d, 7.4 (m, 5H), 7.4 (m		555.4	1.6 (d, J=7.1 Hz, 3H), 2.8 (d,
(qn J=7.0 Hz, 1H), 5.7 (broad, 1H), 6.4 (d, J=5.3 Hz, 1H), 6.7 (broad, 1H), 7.1 (m, 2H), 7.3 (m, 1H), 7.4 (m, 4H), 7.6 (m, 3H), 8.1 (d, J=5.3 Hz, 1H), 8.7 (broad, 1H). 525.4 1.6 (d, J=6.9 Hz, 3H), 2.4 (s, 3H), 3.4 (s, 2H), 3.8 (m, 5H), 5.2 (qn J=6.8 Hz, 1H), 5.8 (broad, 1H), 6.4 (d, J=5.2 Hz, 1H), 6.7 (broad, 1H), 7.1 (m, 2H), 7.3 (m, 1H), 7.4 (m, 4H), 7.6 (m, 3H), 8.1 (d, J=5.2 Hz, 1H), 8.7 (broad, 1H). C006 497.4 2.2 (s, 3H), 2.7 (2.7, 2H), 3.4 (s, 3H), 3.6 (t, J=5.7 Hz, 2H), 3.7 (s, 2H), 4.8 (d, J=5.9 Hz, 2H), 5.8 (broad, 1H), 6.5 (d, J=5.3 Hz, 1H), 6.8 (broad, 1H), 7.1 (m, 2H), 7.4 (m, 5H), 7.6 (m, 3H), 8.1 (d, 7.4 (m, 5H), 7.4	n-√n		J=5.7 Hz, 4H), 3.4 (s, 6H), 3.5 (t,
1H), 6.4 (d, J=5.3 Hz, 1H), 6.7 (broad, 1H), 7.1 (m, 2H), 7.3 (m, 1H), 7.4 (m, 4H), 7.6 (m, 3H), 8.1 (d, J=5.3 Hz, 1H), 8.7 (broad, 1H). 525.4 1.6 (d, J=6.9 Hz, 3H), 2.4 (s, 3H), 3.4 (s, 2H), 3.8 (m, 5H), 5.2 (qn J=6.8 Hz, 1H), 5.8 (broad, 1H), 6.4 (d, J=5.2 Hz, 1H), 6.7 (broad, 1H), 7.1 (m, 2H), 7.3 (m, 1H), 7.4 (m, 4H), 7.6 (m, 3H), 8.1 (d, J=5.2 Hz, 1H), 8.7 (broad, 1H). C006 497.4 2.2 (s, 3H), 2.7 (2.7, 2H), 3.4 (s, 3H), 3.6 (t, J=5.7 Hz, 2H), 3.7 (s, 2H), 4.8 (d, J=5.9 Hz, 2H), 5.8 (broad, 1H), 6.5 (d, J=5.3 Hz, 1H), 6.8 (broad, 1H), 7.1 (m, 2H), 7.4 (m, 5H), 7.6 (m, 3H), 8.1 (d, 1H), 6.8 (broad, 1H), 7.1 (m, 2H), 7.4 (m, 5H), 7.6 (m, 3H), 8.1 (d, 1H), 6.8 (broad, 1H), 7.1 (m, 2H), 7.4 (m, 5H), 7.6 (m, 3H), 8.1 (d, 1H), 6.8 (broad, 1H), 7.1 (m, 2H), 7.4 (m, 5H), 7.6 (m, 3H), 8.1 (d, 1H), 6.8 (broad, 1H), 7.1 (m, 2H), 7.4 (m, 5H), 7.6 (m, 3H), 8.1 (d, 1H), 6.8 (broad, 1H), 7.1 (m, 2H), 7.4 (m, 5H), 7.6 (m, 3H), 8.1 (d, 1H), 6.8 (broad, 1H), 7.1 (m, 2H), 7.4 (m, 5H), 7.6 (m, 3H), 8.1 (d, 1H), 6.8 (broad, 1H), 7.1 (m, 2H), 7.4 (m, 5H), 7.6 (m, 3H), 8.1 (d, 1H), 6.8 (broad, 1H), 7.1 (m, 2H), 7.4 (m, 5H), 7.6 (m, 3H), 8.1 (d, 1H), 6.8 (broad, 1H), 7.1 (m, 2H), 7.4 (m, 5H), 7.6 (m, 3H), 8.1 (d, 1H), 6.8 (broad, 1H), 7.1 (m, 2H), 7.4 (m, 5H), 7.6 (m, 3H), 8.1 (d, 1H), 6.8 (broad, 1H), 7.1 (m, 2H), 7.4 (m, 5H), 7.6 (m, 3H), 8.1 (d, 1H), 6.8 (broad, 1H), 7.1 (m, 2H), 7.4 (m, 5H), 7.6 (m, 3H), 8.1 (d, 1H), 6.8 (broad, 1H), 6.8	() N	ļ	J=5.8 Hz, 4H), 3.8 (s, 2H), 5.2
(broad, 1H), 7.1 (m, 2H), 7.3 (m, 1H), 7.4 (m, 4H), 7.6 (m, 3H), 8.1 (d, J=5.3 Hz, 1H), 8.7 (broad, 1H). 525.4 1.6 (d, J=6.9 Hz, 3H), 2.4 (s, 3H), 3.4 (s, 2H), 3.8 (m, 5H), 5.2 (qn J=6.8 Hz, 1H), 5.8 (broad, 1H), 6.4 (d, J=5.2 Hz, 1H), 6.7 (broad, 1H), 7.1 (m, 2H), 7.3 (m, 1H), 7.4 (m, 4H), 7.6 (m, 3H), 8.1 (d, J=5.2 Hz, 1H), 8.7 (broad, 1H). C006 497.4 2.2 (s, 3H), 2.7 (2.7, 2H), 3.4 (s, 3H), 3.6 (t, J= 5.7 Hz, 2H), 3.7 (s, 2H), 4.8 (d, J=5.9 Hz, 2H), 5.8 (broad, 1H), 6.5 (d, J=5.3 Hz, 1H), 6.8 (broad, 1H), 7.1 (m, 2H), 7.4 (m, 5H), 7.6 (m, 3H), 8.1 (d, 1H), 6.8 (broad, 1H), 7.1 (m, 2H), 7.4 (m, 5H), 7.6 (m, 3H), 8.1 (d, 1H), 6.8 (broad, 1H), 7.1 (m, 2H), 7.4 (m, 5H), 7.6 (m, 3H), 8.1 (d, 1H), 6.1 (d, 1H), 7.1 (m, 2H), 7.4 (m, 5H), 7.6 (m, 3H), 8.1 (d, 1H), 6.1 (d, 1H), 7.1 (m, 2H), 7.4 (m, 5H), 7.6 (m, 3H), 8.1 (d, 1H), 6.1 (d, 1H)	N		(qn J=7.0 Hz, 1H), 5.7 (broad,
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C005 (d, J=5.3 Hz, 1H), 8.7 (broad, 1H). 525.4 1.6 (d, J=6.9 Hz, 3H), 2.4 (s, 3H), 3.4 (s, 2H), 3.8 (m, 5H), 5.2 (qn J=6.8 Hz, 1H), 5.8 (broad, 1H), 6.4 (d, J=5.2 Hz, 1H), 6.7 (broad, 1H), 7.1 (m, 2H), 7.3 (m, 1H), 7.4 (m, 4H), 7.6 (m, 3H), 8.1 (d, J=5.2 Hz, 1H), 8.7 (broad, 1H). C006 497.4 2.2 (s, 3H), 2.7 (2.7, 2H), 3.4 (s, 3H), 3.6 (t, J=5.7 Hz, 2H), 3.7 (s, 2H), 4.8 (d, J=5.9 Hz, 2H), 5.8 (broad, 1H), 6.5 (d, J=5.3 Hz, 1H), 6.8 (broad, 1H), 7.1 (m, 2H), 7.4 (m, 5H), 7.6 (m, 3H), 8.1 (d, 1H), 7.1 (m, 2H), 7.4 (m, 5H), 7.6 (m, 3H), 8.1 (d, 1H), 7.1 (m, 2H), 7.4 (m, 5H), 7.6 (m, 3H), 8.1 (d, 1H), 7.1 (m, 2H), 7.4 (m, 5H), 7.6 (m, 3H), 8.1 (d, 1H), 7.1 (m, 2H), 7.8 (m, 5H), 7.6 (m, 3H), 8.1 (d, 1H), 7.1 (m, 5H), 7.8 (m, 7H), 8.1 (d, 1H), 7.1 (m, 7H), 7.8 (m, 7H), 8.1 (d, 1H), 7.1 (m, 7H), 7.8 (m, 7H), 8.1 (d, 1H), 7.1 (m, 7H), 7.1 (1H), 7.4 (m, 4H), 7.6 (m, 3H), 8.1
C005 IH). 1.6 (d, J=6.9 Hz, 3H), 2.4 (s, 3H), 3.4 (s, 2H), 3.8 (m, 5H), 5.2 (qn J=6.8 Hz, 1H), 5.8 (broad, 1H), 6.4 (d, J=5.2 Hz, 1H), 6.7 (broad, 1H), 7.1 (m, 2H), 7.3 (m, 1H), 7.4 (m, 4H), 7.6 (m, 3H), 8.1 (d, J=5.2 Hz, 1H), 8.7 (broad, 1H). C006 497.4 2.2 (s, 3H), 2.7 (2.7, 2H), 3.4 (s, 3H), 3.6 (t, J=5.7 Hz, 2H), 3.7 (s, 2H), 4.8 (d, J=5.9 Hz, 2H), 5.8 (broad, 1H), 6.5 (d, J=5.3 Hz, 1H), 6.8 (broad, 1H), 7.1 (m, 2H), 7.4 (m, 5H), 7.6 (m, 3H), 8.1 (d,	Ţ	1	(d, J=5.3 Hz, 1H), 8.7 (broad,
3.4 (s, 2H), 3.8 (m, 5H), 5.2 (qn J=6.8 Hz, 1H), 5.8 (broad, 1H), 6.4 (d, J=5.2 Hz, 1H), 6.7 (broad, 1H), 7.1 (m, 2H), 7.3 (m, 1H), 7.4 (m, 4H), 7.6 (m, 3H), 8.1 (d, J=5.2 Hz, 1H), 8.7 (broad, 1H). C006 497.4 2.2 (s, 3H), 2.7 (2.7, 2H), 3.4 (s, 3H), 3.6 (t, J=5.7 Hz, 2H), 3.7 (s, 2H), 4.8 (d, J=5.9 Hz, 2H), 5.8 (broad, 1H), 6.5 (d, J=5.3 Hz, 1H), 6.8 (broad, 1H), 7.1 (m, 2H), 7.4 (m, 5H), 7.6 (m, 3H), 8.1 (d, 1H), 6.8 (broad, 1H), 7.1 (m, 2H), 7.4 (m, 5H), 7.6 (m, 3H), 8.1 (d, 1H), 6.8 (broad, 1H), 7.1 (m, 2H), 7.4 (m, 5H), 7.6 (m, 3H), 8.1 (d, 1H), 6.8 (broad, 1H), 7.1 (m, 2H), 7.4 (m, 5H), 7.6 (m, 3H), 8.1 (d, 1H), 6.8 (broad, 1H), 7.1 (m, 2H), 7.4 (m, 5H), 7.6 (m, 3H), 8.1 (d, 1H), 6.8 (broad, 1H), 7.1 (m, 2H), 7.4 (m, 5H), 7.6 (m, 3H), 8.1 (d, 1H), 6.8 (broad, 1H), 7.1 (m, 2H), 7.4 (m, 5H), 7.6 (m, 3H), 8.1 (d, 1H), 6.8 (broad, 1H), 7.1 (m, 2H), 7.4 (m, 5H), 7.6 (m, 3H), 8.1 (d, 1H), 6.8 (broad, 1H), 7.1 (m, 2H), 7.4 (m, 5H), 7.6 (m, 3H), 8.1 (d, 1H), 6.8 (broad, 1H), 7.1 (m, 2H), 7.4 (m, 5H), 7.6 (m, 3H), 8.1 (d, 1H), 6.8 (broad, 1H), 7.1 (m, 2H), 7.4 (m, 5H), 7.6 (m, 3H), 8.1 (d, 1H), 6.8 (broad, 1H), 7.1 (m, 2H), 7.4 (m, 5H), 7.6 (m, 3H), 8.1 (d, 1H), 6.8 (broad, 1H), 7.1 (m, 2H), 7.4 (m, 5H), 7.6 (m, 3H), 8.1 (d, 1H), 6.8 (broad, 1H), 7.1 (m, 2H), 7.4 (m, 5H), 7.6 (m, 3H), 8.1 (d, 1H), 6.8 (broad, 1H), 7.1 (m, 2H), 7.4 (m, 5H), 7.6 (m, 3H), 8.1 (d, 1H), 6.8 (broad, 1H), 7.1 (m, 2H), 7.4 (m, 5H), 7.6 (m, 3H), 8.1 (d, 1H), 6.8 (broad, 1H), 7.1 (m, 2H), 7.4 (m, 5H), 7.6 (m, 3H), 8.1 (d, 1H), 7.8 (m, 2H), 8.1 (d, 1H), 9.1 (d, 1H), 9	C005		•
3.4 (s, 2H), 3.8 (m, 5H), 5.2 (qn J=6.8 Hz, 1H), 5.8 (broad, 1H), 6.4 (d, J=5.2 Hz, 1H), 6.7 (broad, 1H), 7.1 (m, 2H), 7.3 (m, 1H), 7.4 (m, 4H), 7.6 (m, 3H), 8.1 (d, J=5.2 Hz, 1H), 8.7 (broad, 1H). C006 2.2 (s, 3H), 2.7 (2.7, 2H), 3.4 (s, 3H), 3.6 (t, J=5.7 Hz, 2H), 3.7 (s, 2H), 4.8 (d, J=5.9 Hz, 2H), 5.8 (broad, 1H), 6.5 (d, J=5.3 Hz, 1H), 6.8 (broad, 1H), 7.1 (m, 2H), 7.4 (m, 5H), 7.6 (m, 3H), 8.1 (d, 1H), 6.8 (broad, 1H), 7.1 (m, 2H), 7.4 (m, 5H), 7.6 (m, 3H), 8.1 (d, 1H), 6.8 (broad, 1H), 7.1 (m, 2H), 7.4 (m, 5H), 7.6 (m, 3H), 8.1 (d, 1H), 6.8 (broad, 1H), 7.1 (m, 2H), 7.4 (m, 5H), 7.6 (m, 3H), 8.1 (d, 1H), 6.8 (broad, 1H), 7.1 (m, 2H), 7.4 (m, 5H), 7.6 (m, 3H), 8.1 (d, 1H), 6.8 (broad, 1H), 7.1 (m, 2H), 7.4 (m, 5H), 7.6 (m, 3H), 8.1 (d, 1H), 6.8 (broad, 1H), 7.1 (m, 2H), 7.4 (m, 5H), 7.6 (m, 3H), 8.1 (d, 1H), 6.8 (broad, 1H), 7.1 (m, 2H), 7.4 (m, 5H), 7.6 (m, 3H), 8.1 (d, 1H), 6.8 (broad, 1H), 7.1 (m, 2H), 7.4 (m, 5H), 7.6 (m, 3H), 8.1 (d, 1H), 6.8 (broad, 1H), 7.1 (m, 2H), 7.4 (m, 5H), 7.6 (m, 3H), 8.1 (d, 1H), 6.8 (broad, 1H), 7.1 (m, 2H), 7.4 (m, 5H), 7.6 (m, 3H), 8.1 (d, 1H), 6.8 (broad, 1H), 7.1 (m, 2H), 7.4 (m, 5H), 7.6 (m, 3H), 8.1 (d, 1H), 6.8 (broad, 1H), 7.1 (m, 2H), 7.4 (m, 5H), 7.6 (m, 3H), 8.1 (d, 1H), 6.8 (broad, 1H), 7.1 (m, 2H), 7.4 (m, 5H), 7.6 (m, 3H), 8.1 (d, 1H), 6.8 (broad, 1H), 7.1 (m, 2H), 7.4 (m, 5H), 7.6 (m, 3H), 8.1 (d, 1H), 6.8 (broad, 1H), 7.1 (m, 2H), 7.4 (m, 5H), 7.6 (m, 3H), 8.1 (d, 1H), 6.8 (broad, 1H), 7.1 (m, 2H), 7.4 (m, 5H), 7.6 (m, 3H), 8.1 (d, 1H), 9.1 (d,		525.4	1.6 (d, J=6.9 Hz, 3H), 2.4 (s, 3H).
D=6.8 Hz, 1H), 5.8 (broad, 1H), 6.4 (d, J=5.2 Hz, 1H), 6.7 (broad, 1H), 7.1 (m, 2H), 7.3 (m, 1H), 7.4 (m, 4H), 7.6 (m, 3H), 8.1 (d, J=5.2 Hz, 1H), 8.7 (broad, 1H). C006 497.4 2.2 (s, 3H), 2.7 (2.7, 2H), 3.4 (s, 3H), 3.6 (t, J=5.7 Hz, 2H), 3.7 (s, 2H), 4.8 (d, J=5.9 Hz, 2H), 5.8 (broad, 1H), 6.5 (d, J=5.3 Hz, 1H), 6.8 (broad, 1H), 7.1 (m, 2H), 7.4 (m, 5H), 7.6 (m, 3H), 8.1 (d, 1H), 6.8 (broad, 1H), 7.1 (m, 2H), 7.4 (m, 5H), 7.6 (m, 3H), 8.1 (d, 1H), 8.1	Ę	1	• •
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(m, 4H), 7.6 (m, 3H), 8.1 (d, J=5.2 Hz, 1H), 8.7 (broad, 1H). C006 497.4 2.2 (s, 3H), 2.7 (2.7, 2H), 3.4 (s, 3H), 3.6 (t, J= 5.7 Hz, 2H), 3.7 (s, 2H), 4.8 (d, J=5.9 Hz, 2H), 5.8 (broad, 1H), 6.5 (d, J=5.3 Hz, 1H), 6.8 (broad, 1H), 7.1 (m, 2H), 7.4 (m, 5H), 7.6 (m, 3H), 8.1 (d, 1H), 7.5 (m, 3H), 8.1 (N		
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C006 497.4 2.2 (s, 3H), 2.7 (2.7, 2H), 3.4 (s, 3H), 3.6 (t, J= 5.7 Hz, 2H), 3.7 (s, 2H), 4.8 (d, J=5.9 Hz, 2H), 5.8 (broad, 1H), 6.5 (d, J=5.3 Hz, 1H), 6.8 (broad, 1H), 7.1 (m, 2H), 7.4 (m, 5H), 7.6 (m, 3H), 8.1 (d,			j
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C006 497.4 2.2 (s, 3H), 2.7 (2.7, 2H), 3.4 (s, 3H), 3.6 (t, J= 5.7 Hz, 2H), 3.7 (s, 2H), 4.8 (d, J=5.9 Hz, 2H), 5.8 (broad, 1H), 6.5 (d, J=5.3 Hz, 1H), 6.8 (broad, 1H), 7.1 (m, 2H), 7.4 (m, 5H), 7.6 (m, 3H), 8.1 (d,			
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3H), 3.6 (t, J= 5.7 Hz, 2H), 3.7 (s, 2H), 4.8 (d, J=5.9 Hz, 2H), 5.8 (broad, 1H), 6.5 (d, J=5.3 Hz, 1H), 6.8 (broad, 1H), 7.1 (m, 2H), 7.4 (m, 5H), 7.6 (m, 3H), 8.1 (d,	C006		
3H), 3.6 (t, J= 5.7 Hz, 2H), 3.7 (s, 2H), 4.8 (d, J=5.9 Hz, 2H), 5.8 (broad, 1H), 6.5 (d, J=5.3 Hz, 1H), 6.8 (broad, 1H), 7.1 (m, 2H), 7.4 (m, 5H), 7.6 (m, 3H), 8.1 (d,		497.4	2.2 (s, 3H), 2.7 (2.7, 2H), 3.4 (s,
2H), 4.8 (d, J=5.9 Hz, 2H), 5.8 (broad, 1H), 6.5 (d, J=5.3 Hz, 1H), 6.8 (broad, 1H), 7.1 (m, 2H), 7.4 (m, 5H), 7.6 (m, 3H), 8.1 (d,			
(broad, 1H), 6.5 (d, J=5.3 Hz, 1H), 6.8 (broad, 1H), 7.1 (m, 2H), 7.4 (m, 5H), 7.6 (m, 3H), 8.1 (d,		-	
N N N 1H), 6.8 (broad, 1H), 7.1 (m, 2H), 7.4 (m, 5H), 7.6 (m, 3H), 8.1 (d,	N		ł
7.4 (m, 5H), 7.6 (m, 3H), 8.1 (d,	N N		
	N N-		t .
F O O			
F	N		5 - 1 - 112, 1112, 511 (bload, 1112).
F	-0		·
	F		·
C007	C007		

EXAMPLE	M+1	NMR(CDCl3)
C008	481.5	1.1 (broad, 6H), 2.6 (broad, 4H), 3.6 (s, 2H), 4.8 (d, J=5.9 Hz, 2H), 5.8 (broad, 1H), 6.5 (d, J=5.3 Hz, 1H), 6.8 (broad, 1H), 7.1 (m, 2H), 7.3 (m, 1H), 7.4 (m, 4H), 7.6 (m, 3H), 8.1 (d, J=5.2 Hz, 1H), 9.1 (broad, 1H).
F—NNNNNNNNNNNNNNNNNNNNNNNNNNNNNNNNNNNN	495.4	1.0 (t, J=7.2 Hz, 3H), 1.6 (m, 2H), 1.6 (d, J=6.8 Hz, 3H), 2.2 (s, 3H), 4.4 (t, J=7.2 Hz, 2H), 3.6 (s, 2H), 5.2 (qn J=6.8 Hz, 1H), 5.8 (broad, 1H), 6.4 (d, J=5.3 Hz, 1H), 6.7 (broad, 1H), 7.1 (m, 2H), 7.3 (m, 1H), 7.4 (m, 5H), 7.6 (m, 2H), 8.1 (d, J=5.3 Hz, 1H), 8.7 (broad, 1H).
C010 N N N N N N N N N N N N N N N N N N	524.4	1.6 (d, J=7.0 Hz, 3H), 2.29 (s, 2H), 2.33 (s, 3H), 2.6 (m, 4H), 3.6 (s, 2H), 5.2 (qn J=7.0 Hz, 1H), 5.6 (d, J=6.4 Hz, 1H), 6.4 (d, J=5.3 Hz, 1H), 6.7 (broad, 1H), 7.1 (m, 2H), 7.3 (m, 1H), 7.4 (m, 5H), 7.6 (m, 2H), 8.1 (d, J=5.3 Hz, 1H), 8.7 (broad, 1H).
C011 NH NH N 2	945.8	

EXAMPLE	M+1	NMR(CDCI3)
CO12	538.4	1.6 (d, J=6.8 Hz, 3H), 2.4 (s, 3H), 3.0 (s, 3H), 3.1 (s, 3H), 3.3 (s, 2H), 3.7 (s, 2H), 5.2 (qn J=6.8 Hz, 1H), 5.8 (broad, 1H), 6.4 (d, J=5.4 Hz, 1H), 6.7 (broad, 1H), 7.1 (m, 2H), 7.3 (m, 1H), 7.4 (m, 4H), 7.6 (m, 3H), 8.1 (d, J=5.3 Hz, 1H), 8.7 (broad, 1H).
C013 N N N N N N N N N N N N N N N N N N		1.6 (d, J=6.9 Hz, 3H), 3.4 (s, 2H), 5.2 (qn J=6.9 Hz, 1H), 5.6 (broad, 1H), 6.4 (d, J=5.3 Hz, 1H), 6.6 (broad, 1H), 7.1 (m, 2H), 7.3 (m, 1H), 7.4 (m, 4H), 7.6 (m, 3H), 8.1 (d, J=5.3 Hz, 1H), 8.7 (broad, 1H).
CO14 F N N N N N O N N N N N N N	566.2	1.0 (m, 6H), 1.6 (d, J=6.9 Hz, 3H), 2.4 (s, 3H), 3.3 (s 2H), 3.4 (m, 4H), 3.7 (s, 2H), 5.2 (qn J=6.9 Hz, 1H), 5.7 (broad, 1H), 6.4 (d, J=5.2 Hz, 1H), 6.7 (broad, 1H), 7.1 (m, 2H), 7.3 (m, 1H), 7.4 (m, 4H), 7.6 (m, 3H), 8.1 (d, J=5.2 Hz, 1H), 8.7 (broad, 1H).

EXAMPLE	M+1	NMR(CDCl3)
F N N N N N N N N N N N N N N N N N N N	552.2	1.1 (m, 3H), 1.6 (d, J=6.9 Hz, 3H), 2.4 (2s, 3H), 2.9 & 3.1 (2s 2H), 3.3 (2s, 2H), 3.5 (m, 2H), 3.7 (m, 2H), 5.2 (qn J=6.7 Hz, 1H), 5.7 (broad, 1H), 6.4 (d, J=5.2 Hz, 1H), 6.7 (broad, 1H), 7.1 (m, 2H), 7.3 (m, 1H), 7.4 (m, 4H), 7.6 (m, 3H), 8.1 (d, J=5.2 Hz, 1H), 8.7 (broad, 1H).
C016 N N N N N N N N N N N N N N N N N N N	524.1	1.6 (d, J=6.8 Hz, 3H), 2.4 (s, 3H), 2.9 (d, J=5.1 Hz, 2H), 3.1 (s, 3H), 3.7 (s, 2H), 5.2 (qn J=6.9 Hz, 1H), 5.7 (broad, 1H), 6.4 (d, J=5.3 Hz, 1H), 6.5 (broad, 1H), 7.1 (m, 3H), 7.3 (m, 1H), 7.4 (m, 4H), 7.6 (m, 3H), 8.1 (d, J=5.3 Hz, 1H), 8.7 (broad, 1H).

EXAMPLE	M+1	NMR(CDCl3)
F N N N N N N N N N N N N N N N N N N N	536.2	1.6 (d, J=6.8 Hz, 3H), 2.2 (m, 2H), 2.5 (m, 1H), 2.8 (d, J=5.1 Hz, 3H), 3.5 (m, 1H), 3.6 (m, 1H), 3.8 (m, 2H), 5.2 (qn J=6.9 Hz, 1H), 5.7 (broad, 1H), 6.4 (d, J=5.2 Hz, 1H), 6.5 (broad, 1H), 7.1 (m, 3H), 7.3 (m, 1H), 7.4 (m, 4H), 7.6 (m, 3H), 8.1 (d, J=5.2 Hz, 1H), 8.7 (broad, 1H).
CO18 NH N N N N N N N N N N N N N N N N N N	419.2	1.01(s,9h), 2.46(s, 3H), 3.34(s, 2H), 3.87(s, 2H), 6.32(d, J=5.3 Hz, iH), 7.11(m, 1H), 7.21(m, 2H), 7.60(m, 3H), 8.05(d, J=5.3 Hz, 2H), 9.62(br, 1H),
C019 N N N N N N N N N N N N N N N N N N N	550.2	1.6 (d, J=6.8 Hz, 3H), 2.3 (m, 2H), 2.8 (m, 7H), 3.4 (m, 1H), 3.5 (m, 1H), 4.0 (m, 2H), 5.2 (qn J=7.0 Hz, 1H), 5.7 (broad, 1H), 6.4 (d, J=5.3 Hz, 1H), 6.5 (broad, 1H), 7.1 (m, 3H), 7.3 (m, 1H), 7.4 (m, 5H), 7.6 (m, 2H), 8.1 (d, J=5.3 Hz, 1H), 8.8 (broad, 1H).

5

Ex.	STRUCTURE	ES+ (M+1)
D01	CI N N OH	494.1
D02	CH ₃ OH	406.2
D03	CH ₃	456.2

Ex.	STRUCTURE	ES+
		(M+1)
D04	PNOH	494.2
·	F N N N	
D05	F CH ₃ N—CH ₃	483.3
-	CH ₃	
D06	P OH	416.1 (ES-)
D07	F	454.3
	CH ₃ OH	
	CH ₃	

Ex.	STRUCTURE	ES+
		(M+1)
D08	CH ₃ CH ₃ CH ₃	592.3
	F N N	
D09	CH ₃ OH	440.3
D10	P OH	470.2
D11	CH ₃ CH ₃ CH ₃	433.3

Ex.	STRUCTURE	ES+
		(M+1)
D12	CI	478.2
	N N OH	
D13	P OH	444.2
	F N N N	777.2
	F	
D14	N OH	404.2
	N N N	
н	F	

Ex.	CTDIICTIDE	FC.
EX.	STRUCTURE	ES+
D15	E	(M+1)
D15	N N N N N N N N N N N N N N N N N N N	444.2
	F	
D16	CH ₃ OH	440.2
D17	CI OH	460.3
	F	
D18	CH ₃	420.3
	P OH	

Ex.	STRUCTURE	ES+
	·	(M+1)
D19	ÇH ₃ Ç CH ₃	486.2
	CH ₃ O CH ₃ .	
	N N N	
	ОН	
	F	
D20	CH₃ OH	456.2
	N N	
	F	
D21	F OH	510.2
-	N	
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Ex.	STRUCTURE	ES+
		(M+1)
D22	CH₃	456.2
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	N=	
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}		
	F— N	
	N	
D23	O—CH ₃	486.2
	CH ₃	
	N	
	N≕	
	N N	
	N	
	F—OH	
D24	F	462.3
	F	
	N=\(
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	F—N	
	NOH	

Ex.	STRUCTURE	ES+
DOF		(M+1)
D25		476.3
		1
	N={	
	N N	
	F OH	
D26	CH ₃ HO	486.2
	CH ₃ O	
} .		·
	N N N	
	F	
D27	НО	454.3
	N N	
	CH ₃	
	F	

Ex.	STRUCTURE	ES+
	·	(M+1)
D28	F	462.3
	\ F	
	N	
	N=\	
		,
	Б	
D29		472.2
		.,
	S—CH ₃	! :
	N	
-	N=	
÷	N .	
		-
	F—()—()	•
D30	CH ₃	486.2
1230	\ \rightarrow -0	400.2
	√ O—CH ₃	
	N=	·
	N N	
	F	
	, NOH	

Ex.	CTDIICTIDE	T ==
Ex.	STRUCTURE	ES+
Dat	F. 🐟	(M+1)
D31		376.3
	N N OH	
D32	F	378.2
	N N OH	
	CH ₃ N CH ₃	
D33	Si 13	406.3
	N N OH	
	CH ₃ N	·
D34	F	438.6 (ES-)
	ОН	
	CH ₃	

		
Ex.	STRUCTURE	ES+
		(M+1)
D35	F	444.3
	P N N OH	
D36	F	392.3
	CH ₃ N OH	
D37		406.3
	CH ₃ N OH	·
D38	F	364.3
	N N OH	
	CH ₃ N	

(M+1		CONTRACTOR	т
D39 CH ₃ D41 CH ₃ CH ₃ CH ₃ CH ₃ A54.3	Ex.	STRUCTURE	ES+
D39 CH ₃ D41 CH ₃ CH ₃ CH ₃ CH ₃ A06.3 378.3 406.3 A06.3 A06.3	<u> </u>		(M+1)
D40 CH ₃ N OH 406.3 CH ₃ OH 392.3 CH ₃ CH ₃ A CH ₃ A 454.3	D39		378.3
D40 CH ₃ N OH 406.3 A06.3 CH ₃ N OH 392.3 CH ₃ CH ₃ A CH ₃ A A A A A A A A A A A A A		N	
D40 CH ₃ N OH CH ₃ N OH 392.3 CH ₃			·
D40 CH ₃ N OH CH ₃ N OH 392.3 CH ₃ A 454.3			
D40 CH ₃ OH CH ₃ OH 392.3 CH ₃ OH CH ₃ OH 454.3	1	N OH	
D40 CH ₃ OH CH ₃ OH 392.3 CH ₃ OH CH ₃ OH 454.3	1) T	
D40 CH ₃ OH CH ₃ OH 392.3 CH ₃ OH CH ₃ OH 454.3		CH ₃ N	
D41 F 392.3 CH ₃ N OH CH ₃ N A 454.3	D40	F	406.3
D41			100.0
D41			·
D41	·	N L	
D41		N, N OH	
D41 F N N OH CH ₃ N CH ₃ CH ₃ CH ₃ CH ₃ CH ₃ A A A A A A A A A A A A A			
D41 F N N OH CH ₃ N CH ₃ CH ₃ CH ₃ CH ₃ CH ₃ A A A A A A A A A A A A A		CH ₃ N	
D42 F	D41		392.3
D42 F		N	
D42 F 454.3	-		
D42 F 454.3			
D42 F 454.3		N OH	
D42 F 454.3		Ť	,
CH ₃ (CH ₃ (A54.3)			
CH ₃	D42	F _Y	454.3
		N	
	į		
		CH ₃	
N			
T T		N	
CH ₃		CH ₃	

Γ_	Tama-ramana	T ==
Ex.	STRUCTURE	ES+
ļ		(M+1)
D43	F	468.3
	l N	
	OH OH	
[-	N	
D44	F	408.3
	N	
	N ₂ N OH	
	ÇH₃ ,	
	HONN	
	CH ₃	
D45	F	394.3
	N	
	N OH	
	CH ₃	
ļ	HO N	
D46	F	380.3
]		, , ,
1		
[N N	
	N ₂ N OH	
	HO	

Ex.	STRUCTURE	ES+
		(M+1)
D47	F N	390.3
	N N OH	
	N	
D48	F. N.	394.3
	N N OH	
	CH ₃	
D49		349.2
	N OH	
	HO	
D50	F	408.3
	N N OH	
	CH ₃	

Ex.	CTDICTIDE	EC
EX.	STRUCTURE	ES+
	F. 🐟	(M+1)
D51		406.3
	N N	
1		
	N ₂ N OH	
	N N	1
	, N	
		}
D52	F	462,3
	N	.5_,6
		j L
	F	}
	N N OH	
	T ~	
D53	F	471.3
	N au	
	N CH ₃ N-CH ₃	
·	N N N N TO H ₃	
,	N N	
·		
D54	F	483.3
•		
	CH ₃	
	CH ₃ N-CH ₃	
	N N	
<u> </u>		

Ex.	STRUCTURE	ES+
		(M+1)
D55	FN	467.3
	CH ₃ N-CH ₃	
	N N N	
D56	F ÇH ₃	471.3
	N-CH ₃	
D57	CI N F	505.3
	N—CH ₃	

Ex.	STRUCTURE	ES+
	1 2000	(M+1)
D58	CH ₃	513.3
		•
	N—CH ₃	
D59	P CH ₃	513.4
D33		010.4
	CH ₃	
	N N N-CH ₃	.•
	F CH₃	
D60	S—CH ₃	499.3
	N	
	N—CH ₃	
	F CH ₃	

F.	CODY	
Ex.	STRUCTURE	ES+
<u> </u>		(M+1)
D61	F	497.4
	ÇH₃	
	N-CH ₃	Ì
	N N	}
	Ņ	
1		
D62	F	467.4
		407.4
· .	CH ₃	
	N-CH ₃	
	N ₂ N	
	14	
	, N	
	CH ₃ —	
D63		407.4
		431.4
	; N	
	N N	
	N N	ĺ
	N-CH ₃	
	ĆH₃	
	F	
L		

Ex.	STRUCTURE	ES+
D64	CI	(M+1) 487.3
	N—CH ₃	
<u></u>	F	
D65	CH ₃ CH ₃ N	509.4
	N N N N -CH ₃	
	CH₃	
D66	F. N.	363.5
	ОН	
	CH ₃ N	
	CH₃	

Ex.	STRUCTURE	ES+
	<u> </u>	(M+1)
D67	F.	
100,		417.9
	N	(ES-)
	ОН	
	CH₃ N N	
	CH ₃	ĺ
	CH ₃	
<u> </u>	CH3 CH3	
D68		405.9
	, N	
	OH	
	CH ₃	
	CH ₃ N	
	ĒH₃	
D69	F_	405.7
	NOH	
:		
[[ÇH₃ CH₃	
	CH ₃ N	
	CH ₃	
	Ong	

Ex.	STRUCTURE	ES+
		(M+1)
D70	N OH	517.6
	CH ₃ UH	
D71	N OH	453.8
	CH ₃	
D72	N—CH ₃	446.7
	N CH ₃	
D73	CH ₃ CH ₃ CH ₃	567.7

Ex.	STRUCTURE	ES+
		(M+1)
D74	CH ₃ O CH ₃ CH ₃ CH ₃	553.7
D75	CH ₃ O CH ₃ CH ₃ O CH ₃ CH ₃	537.8
D76	CH ₃ O CH ₃ CH ₃	538.4
D77	CH ₃ O CH ₃ CH ₃ CH ₃ CH ₃	551.7

Ex.	STRUCTURE	ES+
LA.	STRUCTURE	(M+1)
D78	F	466.7
	CH ₃ N—CH ₃	
D79	ĊH ₃	390.7
	CH ₃ CH ₃	
D80	F	404.7
	CH ₃ N-CH ₃	
	CH ₃	

Ex.	STRUCTURE	ES+
		(M+1)
D81	CH ₃ N-CH ₃	432.8
	CH ₃	
D82	CH ₃ N-CH ₃	432.6
D83	CH ₃	402.7

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Ex.	STRUCTURE	ES+
	SIRUCIURE	
D84	F	(M+1) 418.7
	N CH₃	
	CH ₃ CH ₃	
D85	F_	512.7
	CH₃	
	CH ₃	
	CH ₃	70
D86	N N	494.7
	N F	· · · · · ·
	CH ₃ CH ₃	

Ex.	CEDITORIDE	T
EX.	STRUCTURE	ES+
J	<u> </u>	(M+1)
D87	F	432.7
	CH ₃ CH ₃	
·	CH₃ ≟ CH₃	
D88	F_	432.7
	CH ₃ CH ₃ CH ₃	
	CH ₃]
D89	F	434.6
	N CH₃	
	CH ₃ N CH ₃	

Ex.	STRUCTURE	ES+
		(M+1)
D90	F	416.7
	CH ₃	
D91		420.7
	CH ₃ N CH ₃	
D92	F	435.7
	CH ₃ N CH ₃ CH ₃	

Γ		
Ex.	STRUCTURE	ES+
		(M+1)
D93	N=	488.6
	F N F	
	j N	
	· >=/	
	N—CH ₃	
	CH₃	
D94	N	488.6
	N N	
- -	F	·
	F N	
	N	
	CH₃ CH₃	
D95	ÇH₃	400.7
D93) N	480.7
	N	
	CH ₃	
	N	ľ
İ	N	
	<u></u>	•
	cH₃	

Ex.	STRUCTURE	ES+
		(M+1)
D96	CH ₃ O NH ₂	433.7
	ĊH₃ E	
D97	CH ₃ O NH ₂	447.7
D98	E N	475.7
	N CH ₃ O NH ₂	
	CH ₃	<u>.</u>

Ex.	STRUCTURE	ES+
		(M+1)
D99	CH ₃ CH ₃ O NH ₂	523.7
	ĊH ₃	·
D100	N CH ₃ O NH ₂	537.7
D101	CH ₃ O NH ₂	459.9

Ex.	STRUCTURE	ES+
D102	F	(M+1) 475.7
	CH ₃ O NH ₂	
D103	F OH	418.0 (ES-)
D104	F OH	406.5
D105	P CH ₃ CH ₃ CH ₃ CH ₃ OH	420.5

	Ex.	STRUCTURE	ES+
			(M+1)
	D106	CH ₃	452.2
		N	(ES-)
		N NOH	
		N N OH	
۱			
		F	
	D107	ÇH₃ , CH₃	420.5
		N CH ₃	
		N CH ₃ OH	
		· N	,
		- N	-
		F	
	D108		390.4
		N	İ
		И ОН	
ļ	•	N N N	
)—-Ki	
			j
		F	İ
L		Г	

Ex.	STRUCTURE	ES+
		(M+1)
D109	Ņ CH ₃	405.4
	N/CH	
	1	
	CH ₃	
) <u> </u>	j.
	F	
D110	N CH3	419.0
		(ES-)
	N CH ₃	
D111	, CH ₃	401.4
	N N CH₃	
	<u> </u>	

F	COMPATICATE TO THE PARTY OF THE	Τ
Ex.	STRUCTURE	ES+
		(M+1)
D112	H ₂ N	407.4
	N N CH ₃ CH ₃	
D113	NH ₂	007.4
	N CH ₃	387.4
D114	N CH₃	407.4
	N CH ₃ OH	

Ev	CEDICETER	
Ex.	STRUCTURE	ES+
		(M+1)
D115	Ņ CH₃	402.4
	N CH ₃	,
1	N CH ₃	
	F	
D116	Ņ ^{CH} ₃	432.5
	N N CH ₃	
	F	
D117	N CH ₃	420.4
]	CH ₃	
	N N NH ₂	
	N 0	
Ll	F	

Ex.	STRUCTURE	T-0
	SIRUCIURE	ES+
77110	∠CH ₃	(M+1)
D118	CH ₃ CH ₃	459.3
		ES-O
	N A A	
[Ñ N CH³	
1 .		
	CH ₃	
) N	
]]	F	
D119	CH ₃ CH ₃	463.5
	CH₃	400.0
	N	
]]	N N CH ₃ OH	
	»—й	
	()	
		ļ
D120	F ÇH₃	
	>—CH ₃	475.3
	N—/ CH ₃	(ES-)
	,n-<	
	()v	
		·
		•
	м сн _з	
		}
	F CH ₃	

Ex.	STRUCTURE	ES+
·		(M+1)
D121	CH₃	457.5
	N—CH ₃	
	,n-<	
	(<u>)</u>	·
	N CH ₃	
	F CH	
D122	CH ₃ CH ₃	458.5
	N— CH ₃	,
	N—	
	()	
	N CH ₃	
	F	
D123	CH₃	488.5
	CH ₃	
	N—	,
	N	
	N-	
	F CH ₃	<u> </u>

Ev	CEDICOTOR	T
Ex.	STRUCTURE	ES+
		(M+1)
D124	CH ₃	476.5
	N—CH₃ CH₃	
	/ ·	1
	N—	
	N	
	N—CH₃	
	Ň	
,	NH ₂	
	F. d	
D125		405.1
		'
	N CH₃	
	N-CH ₃	
	N	
	<u> </u>	
	CH ₃ /N	
D126	CH ₃	363
	`N I	000
	N	
- }		
	N-CH ₃	
}		
	N	
	F \\\	

F	CAMPLICAMANDE	T
Ex.	STRUCTURE	ES+
	CH	(M+1)
D127	CH ₃	475.8
	ĵ	
· ·	N CH ₃	
1	N	
1.	CH ₃	
	CH₃	
	F SH	
D128	CH ₃	473.7
	N CH₃	
]		
ļ	F ~	
D129	CH ₃	531.8
	CH ₃ N CH ₃ ↓	•
]	N/ NA	
	CH ₃	
	N CH ₃	i.
j	\sim \sim \sim N	
	N O CH ₃	
	F ·	
D130	CH ₃	529.9
	CH3 N	//
	CH₃	
	N CH₃	
	N	
	F' V	

Ex.	STRUCTURE	ES+
ļ		(M+1)
D131		478.8
1		
1		
	, N	
· .	N CH	
	\ \ \ \ \ \ \ \ \ \ \ \ \ \ \ \ \ \ \	
	N—CH ₃	
	N	
D132	ÇH ₃	
מוטן	CH₃	432.8
}	Ņ	
}	N CH₃	
	N-CH ₃	
. !		
7100	F CH₃	
D133		447.8
ŀ	Ń CH³	
	NI O	. ,
	N CH ₃ NH ₂	. [
ľ	N N	
Ì		
	F	

Ex.	STRUCTURE	ES+ (M+1)
D134	CH ₃ CH ₃	418.8
	CH ₃ N—CH ₃	
D135	CH ₃ CH ₃	461.8
	CH ₃ NH ₂	
D136	CH ₃	432.8
	N CH3	
	N-CH ₃	

Ex.	STRUCTURE	ES+
		(M+1)
D137	CH ₃	432.8
	CH ₃ CH ₃ N—CH ₃	
	F	
D138	CH ₃ ONN OU	512.9
	CH ₃ CH ₃ N—CH ₃	
	F	
D139	CH₃ /	446.8
	CH ₃	
	N CH ₃	
	F	
D140	□ N	416.8
	N CH ₃	
	F	

Ex.	STRUCTURE	ES+ (M+1)
D141	НО	431.7
	CH ₃ O N N N F	
D142	но	443.7
	CH ₃ CH ₃ N N N F	
D143	НО	479.7
	CH ₃	

Ex.	STRUCTURE	ES+
		(M+1)
D144	CH ₃ CH ₃	457.8
	CH ₃	·
	ОН	
	F	
D145	HO	443.7
	CH ₃	
	CH ₃	
D146	НО	460.7
	CH ₃ N N N N F	

Ex.	STRUCTURE	ES+ (M+1)
D147	НО	471.7
	N N N N F	
D148	но	465.7
-	N N N N N N N N N N N N N N N N N N N	
D149	но	440.6
	N-N-N-F	

Ex.	STRUCTURE	ES+ (M+1)
D150	НО	446.7
	S N N F	
D151	НО	430.7
	CH ₃	
	N N N N N N N N N N N N N N N N N N N	
D152	но	482.7
·		
	N N N N F	

5

Ex.	STRUCTURE	ES+ (M+1)
D153	HO N N N N N N N N N N N N N N N N N N N	415.8

EXAMPLE E1 to **EXAMPLE E192**elow were made by procedures similar to those described above.

EX. E1

EX. E2

EX. E3

EX. E4

EX. E5

EX. E6

EX. E7

EX. E8

EX. E9

EX. E10	EX. E11	EX. E12
EX. E13	EX. E14	EX. E15
EX. E16	EX. E17	EX. E18
EX. E19	EX. E20	EX. E21
EX. E22	EX. E23	EX. E24

	,	
EX. E25	EX. E26	EX. E27
EX. E28	EX. E29	EX. E30
EX. E31	EX. E32	EX. E33
EX. E34	EX. E35	EX. E36

EX. E37	EX. E38	EX. E39
EX. E40	EX. E41	EX. E42
EX. E43	EX. E44	EX. E45
EX. E46	EX. E47	EX. E48
	HO CH	E TO THE TOTAL PROPERTY OF THE TOTAL PROPERT
EX. E49	EX. E50	EX. E51

EX. E52	EX. E53	EX. E54
EX. E55	EX. E56	EX. E57
EX. E58	EX. E59	EX. E60
EX. E61	EX. E62	EX. E63
EX. E64	EX. E65	EX. E66

EX. E67	EX. E68	EX. E69
EX. E70	EX. E71	EX. E72
	No.	¥ 7 7 7 7 7 7 7 7 7 7 7 7 7 7 7 7 7 7 7
EX. E73	EX. E74	EX. E75
		Z Z Z Z Z Z Z Z Z Z Z Z Z Z Z Z Z Z Z
EX. E76	EX. E77	EX. E78

EX. E79	EX. E80	EX. E81
EX. E82	EX. E83	EX. E84
EX. E85	EX. E86	EX. E87
	## ## ## ## ## ## ## ## ## ## ## ## ##	
EX. E88	EX. E89	EX. E90
30-		
EX. E91	EX. E92	EX. E03
EX. E94	EX. E95	EX. E96

EX. E97	EX. E98	EX. E99
EX. E100	EX. E101	EX. E102
EX. E103	EX. E104	EX. E105
EX. E106	EX. E107	EX. E108
EX. E109	EX. E110	EX. E111

EX. E112	EX. E113	EX. E114
EX. E115	EX. E116	EX. E117
EX. E118	EX. E119	EX. E120
		50000
EX. E121	EX. E122	EX. E123
EX. E124	EX. E125	EX. E126

		
EX. E127	EX. E128	EX. E129
EX. E130	EX. E131	EX. E132
To the text of the		
EX. E133	EX. E134	EX. E135
EX. E136	EX. E137	EX. E138
EX. E139	EX. E140	EX. E141

EX. E142	EX. E143	EX. E144
EX. E145	EX. E146	EX. E147
EX. E148	EX. E149	EX. E150
EX. E151	EX. E152	EX. E153
EX. E154	EX. E155	EX. E156
EX. E157	EX. E158	EX. E159

EX. E160	EX. E161	EX. E162
EX. E163	EX. E164	EX. E165
EX. E166	EX. E167	EX. E168
EX. E169	EX. E170	EX. E171

EX. E172	EX. E173	EX. E174
EX. E175	EX. E176	EX. E177
EX. E178	EX. E179	EX. E180
		¥
EX. E181	EX. E182	EX. E183

EX. E184	EX. E185	EX. E186
EX. E187	EX. E188	EX. E189
EX. E190	EX. E191	EX. E192

EXAMPLE F1 to **EXAMPLE F182** below were made by procedures similar to those described above.

EX. F1	EX. F2	EX. F3
24 2 2 2 2 2 2 2 2 2 2 2 2 2 2 2 2 2 2	2 - 2 - 2 - 2 - 2 - 2 - 2 - 2 - 2 - 2 -	2= 2 2 2 2 2 2 2 2 2 2 2 2 2 2 2 2 2 2

EX. F4	EX. F5	EX. F6
		HAY OH
EX. F7	EX. F8	EX. F9
HN Z Z Z Z Z Z Z Z Z Z Z Z Z Z Z Z Z Z Z	₹	8 Z Z Z Z Z Z Z Z Z Z Z Z Z Z Z Z Z Z Z
EX. F10	EX. F11	EX. F12
EX. F13	EX. F14	EX. F15
HN OH		

EX. F16	EX. F17	EX. F18
EX. F19	EX. F20	EX. F21
EX. F22	EX. F23	EX. F24
EX. F25	EX. F26	EX. F27

EX. F28	EX. F29	EX. F29
EX. F30	EX. F31	EX. F32
HN NI	A A A A A A A A A A A A A A A A A A A	HY NH2
EX. F33	EX. F34	EX. F35
EN CH		
EX. F36	EX. F37	EX. F38

EX. F39	EX. F40	EX. F41
		F N N N N N N N N N N N N N N N N N N N
EX. F42	EX. F43	EX. F44
F NH		F N N N N N N N N N N N N N N N N N N N
EX. F45	EX. F46	EX. F47
	F N N N N N N N N N N N N N N N N N N N	
EX. F48	EX. F49	EX. F50
	NAT NO NO NO NO NO NO NO NO NO NO NO NO NO	NH, NH, NH, NH, NH, NH, NH, NH, NH, NH,

EX. F51	EX. F52	EX. F53
	NH NH NH NH NH NH NH NH NH NH NH NH NH N	
EX. F54	EX. F55	EX. F56
	X	
EX. F57	EX. F58	EX. F59
EX. F60	EX. F61	EX. F62

EX. F63	EX. F64	EX. F65
EX. F66	EX. F67	EX. F68
EX. F69	EX. F70	EX. F71
		B 2 2 2 2 2 2 2 2 2 2 2 2 2 2 2 2 2 2 2
EX. F72	EX. F73	EX. F74
		H N N N N N N N N N N N N N N N N N N N

EX. F75	EX. F78	EX. F79
		Kiriot III
EX. F80	EX. F81	EX. F82
NH4		F Z Z Z
EX. F83	EX. F84	EX. F85
EX. F86	EX. F87	EX. F88

EX. F89	EX. F90	EX. F91
F N N N N N N N N N N N N N N N N N N N		
EX. F92	EX. F93	EX. F94
F N N N N N N N N N N N N N N N N N N N		No.
EX. F95	EX. F96	EX. F97
NH N NH		NH ₂
EX. F98	EX. F99	EX. F100
NH ₂	NH2 N N	No. 1

EX. F101	EX. F102	EX. F103
NA NA NA NA NA NA NA NA NA NA NA NA NA N	B D D D D D D D D D D D D D D D D D D D	NH ₂ NH ₂ NH ₂
EX. F104	EX. F105	EX. F106
CH CH		NH ₂ NH ₂ NH ₂
EX. F107	EX. F108	EX. F109
CH CH	P N N	
EX. F110	EX. F111	EX. F112
OH OH	OH OH	NH ₂
EX. F113	EX. F114	EX. F115

EX. F116	EX. F117	EV 12110
EA. FIIO	EA. FII/	EX. F118
O NH2	NH ₂ OH	NH2 N N OH
EX. F119	EX. F120	EX. F121
NH2 N N N	2 2 2 2 2 2 2 2 2 2 2 2 2 2 2 2 2 2 2	NH2 NH2 NH2
EX. F122	EX. F123	EX. F124
NH ₂	NA NA NA NA NA NA NA NA NA NA NA NA NA N	NA NA NA NA NA NA NA NA NA NA NA NA NA N
EX. F125	EX. F126	
		NH2 NH2

EX. F128	EX. F129	EX. F130
NH2 NH2 NH2 NH2 NH2 NH2 NH2 NH2 NH2 NH2	NH ₂	
EX. F131	EX. F132	EX. F133
EX. F134	EX. F135	EX. F136
	H ₂ N N N N N N N N N N N N N N N N N N N	NH ₂
EX. F137	EX. F138	EX. F139
N N N N N N N N N N N N N N N N N N N	NH ₂	

EX. F140	EX. F141	EX. F142
	NH2 N N N N N N N N N N N N N N N N N N	
EX. F143	EX. F144	EX. F145
EX. F146	EX. F147	EX. F148
N N N N N N N N N N N N N N N N N N N		H ₂ N H ₂ N H ₁ N H ₂ N H ₁ N H ₂ N H ₂ N H ₃ N H ₄ N H ₁ N H ₂ N H ₃ N H ₄ N H ₄ N H ₅ N
EX. F149	EX. F150	EX. F151

EX. F152	EX. F153	EX. F154
NH2 NH2 OH	NH ₂ OH	Note to the second seco
EX. F155	EX. F156	EX. F157
	H-O	PA NO PA NO
EX. F158	EX. F159	EX. F160
Q N N N OH	a N N N OH	HAN HO
EX. F161	EX. F162	EX. F163
Han N N N N N N N N N N N N N N N N N N N		

EX. F164	EX. F165	EX. F166
EX. F167	EX. F168	EX. F169
The state of the s		
EX. F170	EX. F171	EX. F172
	H a H a	
EX. F173	EX. F174	EX. F175

EX. F176	EX. F177	EN 19160
EA. F1/0	EA. F1//	EX. F178
	Ha Ha	
EX. F179	EX. F180	EX. F181
EX. F182		

WHAT IS CLAIMED IS:

1. A compound of the formula (I):

(I)

or a pharmaceutically acceptable salt or hydrate thereof, wherein FusedHet is

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R<sup>1</sup> is H,
                                    -C<sub>1</sub>-6alkyl,
                                   -C(O)(C_{1-6}alkyl),
  5
                                   -C(O)-C_{1-6}alkyl-aryl,
                                   -C<sub>0-4</sub>alkyl-aryl,
                                   -C<sub>0-4</sub>alkyl-indanyl,
                                   -C<sub>0-4</sub>alkyl-imidazolyl,
                                   -C<sub>0-4</sub>alkyl-thiazolyl,
10
                                   -C<sub>0-4</sub>alkyl-pyrazolyl,
                                   -C0_4alkyl-oxadiazolyl,
                                   -C<sub>0-4</sub>alkyl-C<sub>3-6</sub>cycloalkyl,
                                   -C<sub>0</sub>-4alkyl-C<sub>1</sub>-4alkoxy,
                                   -C_1-4alkyl-N(C<sub>0</sub>-4alkyl)(-C<sub>0</sub>-4alkyl),
15
                                   -C1-4alkyl-N(-C0-4alkyl)-CO-C1-4alkoxy,
                                   -C1-4alkyl-piperadinyl,
                                   -C<sub>0-4</sub>alkyl-triazolyl,
                                   -C<sub>1-4</sub>alkyl-imidazothiazolyl,
                                   -C<sub>1-4</sub>alkyl-benzimidazolyl,
20
                                  -C<sub>1-4</sub>alkyl-benzothiazolyl,
                                   -C1-4alkyl-benzotetrahydrofuranyl,
                                  -C<sub>1</sub>-4alkyl-benzodioxolyl,
                                   -C1-4alkyl-(heterocycloC4O2alkyl),
                                   -C1-4alkyl-(heterocycloC5O1alkyl),
25
                                  -C1_4alkyl-tetrahydrofuran, or
                                  -C<sub>1-4</sub>alkyl-oxetanyl;
                            R^{11} is H or -C_{1-6}alkyl;
                           or R1 and R11, together with the N to which they are attached, form
       a morpholinyl;
                           R2, R21, R22 each independently is H, halogen, or -C1-4alkyl;
30
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R<sup>3</sup> is H,
                                     -C<sub>1</sub>-4alkyl,
                                     -C3-6cycloalkyl,
                                     -C<sub>1-4</sub>alkyl-aryl,
 5
                                     -C1_4alkyl-azetidinyl,
                                     -C1_4alkyl-azetidinyl-CO-C0_4alkyl-N(C0_4alkyl)(C0_4alkyl),
                                     -C<sub>1-4</sub>alkyl-pyrrolidinyl,
                                     -C1_4alkyl-piperidinyl,
                                     -C<sub>1-4</sub>alkyl-morpholinyl,
10
                                     -C<sub>0-4</sub>alkyl-N(C<sub>0-4</sub>alkyl)(C<sub>0-4</sub>alkyl),
                                     -C0-4alkyl-N(C0-4alkyl)(C0-4alkyl-C1-4alkoxy),
                                     -C_0-4alkyl-N(C<sub>0</sub>-4alkyl-C<sub>1</sub>-4alkoxy)(C<sub>0</sub>-4alkyl-C<sub>1</sub>-4alkoxy),
                                     -C_{1-4}alkyl-N(C_{0-4}alkyl)-(C_{1-4}alkyl)-aryl,
                                     -C1-4alkyl-N(C0-4alkyl)-C1-4alkyl-tetrahydrofuranyl,
15
                                     -C1_4alkyl-N(C0_4alkyl)-C1_4alkyl-azetidinyl,
                                     -C_{1-4}alkyl-N(C<sub>0-4</sub>alkyl)-C<sub>1-4</sub>alkyl-N(C<sub>0-4</sub>alkyl)(C<sub>0-4</sub>alkyl),
                                     -C_{1-4}alkyl-N(C_{0-4}alkyl)-C_{1-4}alkyl-N(C_{0-4}alkyl)-C_{0-4}alkyl-
        SO<sub>2</sub>C<sub>1-4</sub>alkyl),
                                     -CO-N(C<sub>0-4</sub>alkyl)-C<sub>1-4</sub>alkyl-aryl,
20
                                     -CO-N(C_{0-4}alkyl)-C_{1-4}alkyl-N(C_{0-4}alkyl)(C_{0-4}alkyl),
                                     -C<sub>0</sub>-4alkyl-CO-C<sub>0</sub>-4alkyl,
                                     -C<sub>0</sub>-4alkyl-CO-C<sub>0</sub>-4alkoxy,
                                     -C0-4alkyl-CO-N(C0-4alkyl)-C1-4alkyl-C1-4alkoxy,
                                     -C<sub>0</sub>-4alkyl-CO-N(C<sub>0</sub>-4alkyl)-C<sub>1</sub>-4alkyl-aryl,
25
                                     -C0-4alkyl-CO-piperidinyl,
                                    -C1-4alkyl-N(C0-4alkyl)-C0-4alkyl-CO-C0-4alkyl-N(C0-
       4alkyl)(C<sub>0-4</sub>alkyl),
                                    -C_0-4alkyl-CO-C<sub>0</sub>-4alkyl-N(C<sub>0</sub>-4alkyl)(C<sub>0</sub>-4alkyl),
                                     -O-C<sub>1-4</sub>alkyl-aryl,
30
                                     -C<sub>1</sub>-4alkyl-O-C<sub>1</sub>-4alkyl,
                                     -C_{0-4}alkyl-N(C_{0-4}alkyl)-C_{0-4}alkyl-C_{0-4}alkyl,
                                     -C_{0-4}alkyl-N(C_{0-4}alkyl)-C_{0-4}alkyl-C_{0-4}alkoxy,
                                     -C_0-4alkyl-N(C_0-4alkyl)-C_0-4alkyl-C_0-4alkyl-aryl,
                                     -C<sub>0-4</sub>alkyl-N(C<sub>0-4</sub>alkyl)-C<sub>0-4</sub>alkyl-CO-C<sub>0-4</sub>alkyl(aryl)<sub>2</sub>,
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-C0-4alkyl-N(C0-4alkyl)-C0-4alkyl-CO-C0-4alkyl-pyrrolyl,
                                   -C0-4alkyl-N(C0-4alkyl)-C0-4alkyl-CO-C0-4alkyl-
       pyrrolidinyl,
                                   -C0-4alkyl-N(C0-4alkyl)-C0-4alkyl-CO-C0-4alkyl-
       azetidinyl,
                                   -C0-4alkyl-N(C0-4alkyl)-C0-4alkyl-CO-C2-4alkenyl-
       pyrrolidinyl,
                                  -C0-4alkyl-N(C0-4alkyl)-C0-4alkyl-CO-C0-4alkyl-
       thiophenyl,
10
                                  -C<sub>0-4</sub>alkyl-N(C<sub>0-4</sub>alkyl)-C<sub>0-4</sub>alkyl-C<sub>0-4</sub>alkenyl-
       thiophenyl,
                                   -C0-4alkyl-N(C0-4alkyl)-C0-4alkyl-CO-S-C1-4alkyl-aryl,
                                   -C<sub>0-4</sub>alkyl-N(C<sub>0-4</sub>alkyl)-C<sub>0-4</sub>alkyl-CO-C<sub>3-6</sub>cyclolkyl,
                                   -C<sub>0-4</sub>alkyl-N(C<sub>0-4</sub>alkyl)-C<sub>0-4</sub>alkyl-CO-O-C<sub>1-4</sub>alkyl-aryl,
15
                                   -C<sub>0-4</sub>alkyl-CO-N(C<sub>0-4</sub>alkyl)-C<sub>0-4</sub>alkyl-C<sub>1-4</sub>alkoxy,
                                   -C_1_4alkyl-N(C<sub>0</sub>_4alkyl)(-SO<sub>2</sub>C<sub>1</sub>_4alkyl),
                                  -C_0-4alkyl-N(C<sub>0</sub>-4alkyl)-C<sub>1</sub>-4alkyl-SO<sub>2</sub>C<sub>1</sub>-4alkyl,
                                  -C<sub>0-4</sub>alkyl-S-C<sub>1-4</sub>alkyl-aryl,
                                  -C1-4alkyl-PO(C1-4alkoxy)(C1-4alkoxy),
20
                                  -C1-4alkyl-azetidinyl-CO-N(C0-4alkyl)(C0-4alkyl),
                                  -C<sub>1</sub>-4alkyl-(heterocycloC<sub>4</sub>N<sub>1</sub>O<sub>1</sub>alkyl),
                                  -C<sub>0-4</sub>alkyl-CO-(heterocycloC<sub>5</sub>N<sub>1</sub>alkyl),
                                  -C<sub>0</sub>-4alkyl-CO-N(C<sub>0</sub>-4alkyl)-(heterocycloC<sub>5</sub>N<sub>1</sub>alkyl),
                                  -C1_4alkyl-(heterocycloC4N2alkyl)-C1_4alkyl,
25
                                  -C1_4alkyl-(heterocycloC4N2alkyl)-CO-C0_4alkoxy,
                                  -C1-4alkyl-(heterocycloC4N2alkyl)--C1-4alkyl-N(C0-
       4alkyl)(C<sub>0-4</sub>alkyl),
                                  -C1-4alkyl-(heterobicycloC5N2alkyl)-C1-4alkyl, or
                                  -C1_4alkyl-NH-(heterobicycloC7N1alkyl); and
30
                           R4 is -C1-6alkyl;
                         wherein any of the above aryl, hetaryl, cycloalkyl, or heterocycloalkyl
       optionally is substituted with 1-4 substituents, each substituent independently is
```

halogen, NO2, -CN, -C1_4alkyl, -C0_4alkoxy, -S-C1_4alkyl, or -C0_4alkyl-(CO)-

C₀-4alkoxy; and any of the above alkyl optionally is substituted with 1-4 substituents, each substituent independently is halogen, -N₃, -CN, -COOH, or -C₀-4alkoxy.

2. The compound according to claim 1, wherein FusedHet is

5

or a pharmaceutically acceptable addition salt and/or hydrate thereof.

3. The compound according to claim 1, wherein FusedHet is

10

or a pharmaceutically acceptable addition salt and/or hydrate thereof.

4. The compound according to claim 1, wherein FusedHet is

- or a pharmaceutically acceptable addition salt and/or hydrate thereof.
 - 5. The compound according to claim 1, wherein FusedHet is

or a pharmaceutically acceptable addition salt and/or hydrate thereof.

6. The compound according to claim 1, wherein FusedHet is

or a pharmaceutically acceptable addition salt and/or hydrate thereof.

7. The compound according to claim 1, wherein FusedHet is

- 10 or a pharmaceutically acceptable addition salt and/or hydrate thereof.
 - 8. The compound according to claim 1, wherein FusedHet is

or a pharmaceutically acceptable addition salt and/or hydrate thereof.

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9. The compound according to claim 1, wherein FusedHet is

or a pharmaceutically acceptable addition salt and/or hydrate thereof.

10. The compound according to claim 1, wherein FusedHet is

or a pharmaceutically acceptable addition salt and/or hydrate thereof.

10

5

11. The compound according to claim 1, wherein FusedHet is

or a pharmaceutically acceptable addition salt and/or hydrate thereof.

12. The compound according to Claim 1 represented by

WO 03/000682 PCT/US02/19507

E N N N N N N N N N N N N N N N N N N N	HN-N-N-N-N-N-N-N-N-N-N-N-N-N-N-N-N-N-N-	CF ₃ N 3HCI N NH ₂
CF ₃	CF ₃ CH ₃	CF ₃
CF ₃	CF ₃	CF ₃ N N N CO ₂ CH ₃

CF ₉	CF ₃	CF ₃
CF ₃ NH ₂ NH ₂	CF ₃ NHSO₂CH ₃	CF ₃ CO ₂ H CO ₂ H
CF ₃	CF ₃	CF ₃

13. The compound according to Claim 1 represented by:

5 wherein Ar, R, and Z are

Ar Group	R Group	Z Group
2,4-Difluorophenyl	NHCH ₂ C(CH ₃) ₃	CH₂OH
2,4-Difluorophenyl	NHCH ₂ C(CH ₃) ₃	CH ₂ N(CH ₃) ₂
2,4-Difluorophenyl	HN	CH₂OH
2,4-Difluorophenyl	HN	CH ₂ N(CH ₃) ₂
2,4-Difluorophenyl	NHCH ₂ C(CH ₃) ₃	CH₂OCH₃
3-Trifluoromethylphenyl	NHCH₂C(CH₃)₃	CH ₂ -NO
3-Trifluoromethylphenyl	NHCH ₂ C(CH ₃) ₃	CH ₂ -N N-NMe ₂
3-Trifluoromethylphenyl	NHCH₂C(CH₃)₃	CH ₂ -NN-
3-Trifluoromethylphenyl	NHCH ₂ C(CH ₃) ₃	CH₂OH
3-Trifluoromethylphenyl	NHCH2C(CH3)3	CH ₂ N(CH ₃) ₂

		
Ar Group	R Group	Z Group
2-Chloro-4-fluorophenyl	NHCH ₂ C(CH ₃) ₃	СН₂ОН
2-Chloro-4-fluorophenyl	NHCH ₂ C(CH ₃) ₃	СНО
2-Chloro-4-fluorophenyl	NHCH ₂ C(CH ₃) ₃	CH ₂ N(CH ₃) ₂
2-Chlorophenyl	NHCH2C(CH3)3	CH ₂ OH
2-Chlorophenyl	NHCH ₂ C(CH ₃) ₃	CH ₂ N(CH ₃) ₂
4-Chlorophenyl	NHCH ₂ C(CH ₃) ₃	CH₂OH
4-Chlorophenyl	NHCH ₂ C(CH ₃) ₃	CH ₂ N(CH ₃) ₂
3,4-Dichlorophenyl	NHCH2C(CH3)3	СН ₂ ОН
3,4-Dichlorophenyl	NHCH2C(CH3)3	CH ₂ N(CH ₃) ₂
2,3-Dichlorophenyl	NHCH ₂ C(CH ₃) ₃	CH ₂ OH
2,3-Dichlorophenyl	NHCH ₂ C(CH ₃) ₃	CH ₂ N(CH ₃) ₂
4-Fluorophenyl	NHCH ₂ C(CH ₃) ₃	Н
4-Fluorophenyl	NHCH ₂ C(CH ₃) ₂ CH ₂ OH	н
4-Fluorophenyl	NHCH2C(CH3)2CH2OH	CH₂NHCH₃
4-Fluorophenyl	NHCH ₂ C(CH ₃) ₂ CH ₂ OH	CH ₂ N(CH ₃)SO ₂ CH
		3 .
4-Fluorophenyl	HN	СН3
	(R)	
4-Fluorophenyl	NHCH₂C(CH₃)₃	CH₂NHSO₂CH₃
4-Fluorophenyl	NHCH₂C(CH₃)₃	CH₂N(CH₃)SO₂CH
		3
4-Fluorophenyl	NHCH ₂ C(CH ₃) ₃	CH ₂ PO(OMe) ₂
4-Fluorophenyl	NHCH ₂ C(CH ₃) ₃	CH ₂ SO ₂ CH ₃
4-Fluorophenyl	NHCH ₂ C(CH ₃) ₃	СНО

wherein Ar and R are:

Ar Group	R Group
2,4-Difluorophenyl	NHCH ₂ C(CH ₃) ₃
3-Trifluoromethylphenyl	NHCH ₂ C(CH ₃) ₃
3-Trifluoromethylphenyl	(S) NH
3-Trifluoromethylphenyl	NH(CH ₂) ₃ OCH ₃
4-Fluorophenyl	CI
4-Fluorophenyl	F ₃ C NH
4-Fluorophenyl	HN OH (R,S)
4-Fluorophenyl	NHCH2C(CH3)2CH2OH
4-Fluorophenyl	(S) NH
4-Fluorophenyl	(S) NH
4-Fluorophenyl	(S) NH
4-Fluorophenyl	NH(CH ₂) ₃ OCH ₃
4-Fluorophenyl	NH

Ar Group	R Group
4-Fluorophenyl	HN
4-Fluorophenyl	H ₉ C NH
4-Fluorophenyl	HN OH (R)
4-Fluorophenyl	F ₃ C S
4-Fluorophenyl	NH CO ₂ H
4-Fluorophenyl	HN OH (S)
4-Fluorophenyl	OH (R)
4-Fluorophenyl	OH OH (S,S)
4-Fluorophenyl	MeS
4-Fluorophenyl	HN OMe (R)
4-Fluorophenyl	HN (R,S)

Ar Group	R Group
4-Fluorophenyl	HN (R,S)
4-Fluorophenyl	HN

15. The compound according to Claim 1 represented by:

R Group	Z Group
NHCH ₂ C(CH ₃) ₃	CON(OMe)Me
NHCH ₂ C(CH ₃) ₃	СНО
NHCH₂C(CH₃)₃	CH₂OH
NHCH ₂ C(CH ₃) ₃	CH ₂ N(CH ₃) ₂
HN (S)	CON(OMe)Me
HN (S)	СНО
HN (S)	CH ₂ OH
HN (S)	CH ₂ N(CH ₃) ₂

5 or a pharmaceutically acceptable salt thereof.

16. The compound according to Claim 1 represented by:

wherein Ar, R, and Z are

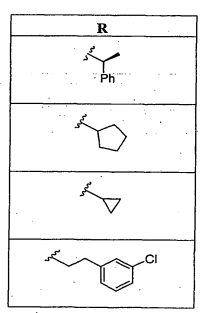
5

	- 	
Ar Group	R Group	Z Group
4-Fluorophenyl	NHCH ₂ C(CH ₃) ₃	CH ₂ OH
4-Fluorophenyl	NHCH ₂ C(CH ₃) ₃	CON(OMe)Me
3-Trifluoromethylphenyl	NHCH ₂ C(CH ₃) ₃	CON(OMe)Me
3-Trifluoromethylphenyl	NHCH ₂ C(CH ₃) ₃	CH ₂ N(CH ₃) ₂
2-Chlorophenyl	NHCH ₂ C(CH ₃) ₃	CH ₂ N(CH ₃) ₂
2-Chloro-4-fluorophenyl	NHCH ₂ C(CH ₃) ₃	CH ₂ N(CH ₃) ₂
2,4-Difluorophenyl	NHCH ₂ C(CH ₃) ₃	CH ₂ N(CH ₃) ₂
2,4-Difluorophenyl	NHCH ₂ C(CH ₃) ₃	CH ₂ OH
2,4-Difluorophenyl	NHCH ₂ C(CH ₃) ₃	CON(OMe)Me
2,4-Difluorophenyl	NHCH ₂ C(CH ₃) ₃	CH ₂ -N N-
2,4-Difluorophenyl	NHCH ₂ C(CH ₃) ₃	CH ₂ -N NNMe ₂
2,4-Difluorophenyl	NHCH ₂ C(CH ₃) ₃	CH ₂ -N NH
2,4-Difluorophenyl	NHCH ₂ C(CH ₃) ₃	CH ₂ -NO
2,4-Difluorophenyl	NHCH ₂ C(CH ₃) ₃	CH ₂ -NH
2,4-Difluorophenyl	NHCH₂C(CH₃)₃	CH ₂ NH(CH ₂) ₂ OCH ₃
2,4-Difluorophenyl	NHCH ₂ C(CH ₃) ₃	CH ₂ NH(CH ₂) ₂ N(CH ₃
2,4-Difluorophenyl	NH(CH₂)₃OCH₃	CON(OMe)Me

		~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~
Ar Group	R Group	Z Group
2,4-Difluorophenyl	H ₃ C NH	CON(OMe)Me
2,4-Difluorophenyl	HN OH (R)	CON(OMe)Me
2,4-Difluorophenyl	NHCH ₂ C(CH ₃) ₂ C H ₂ OH	CH₂OH
2,4-Difluorophenyl	HN	CON(OMe)Me
2,4-Difluorophenyl	HN (S)	CH ₂ N(CH ₃) ₂
2,4-Difluorophenyl	HN (S)	CH₂-NO
2,4-Difluorophenyl	HN SS	CH₂-N NH
2,4-Difluorophenyl	NHCH ₂ C(CH ₃) ₂ C H ₂ OH	CH ₂ N(CH ₃) ₂
2,4-Difluorophenyl	NH(CH ₂) ₃ OCH ₃	CH ₂ -NO
2,4-Difluorophenyl	NH(CH ₂) ₃ OCH ₃	CH ₂ N(CH ₃) ₂
2,4-Difluorophenyl	NHCH₂C(CH₃)₂C H₂OH	CON(OMe)Me
2,4-Difluorophenyl	NH(CH ₂) ₄ OH	CH ₂ N(CH ₃) ₂
2,4-Difluorophenyl	H³C HN	CH ₂ N(CH ₃) ₂
2,4-Difluorophenyl	(E) (D)	CH ₂ N(CH ₃) ₂

		T
Ar Group	R Group	Z Group
2,4-Difluorophenyl	HN	CH ₂ N(CH ₃) ₂
2,4-Difluorophenyl	HN	CH ₂ N(CH ₃) ₂
2,4-Difluorophenyl	HN (S)	CH₂N(CH₃)₂
2,4-Difluorophenyl	HN (S)	CH ₂ N(CH ₃) ₂
2,4-Difluorophenyl	HN (S)	CH ₂ N(CH ₃) ₂
2,4-Difluorophenyl	HN (S)	CH₂N(CH₃)₂
2,4-Difluorophenyl	HN (S)	CH ₂ N(CH ₃) ₂
	CN	
2,4-Difluorophenyl	HN (S)	CH ₂ N(CH ₃) ₂
2,4-Difluorophenyl	(S) OMe	CH ₂ N(CH ₃) ₂
2,4-Difluorophenyl	NH(CH ₂) ₃ CO ₂ H	CH ₂ N(CH ₃) ₂

wherein R is



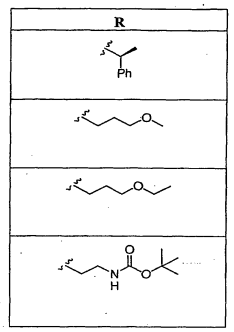
or a pharmaceutically acceptable salt thereof.

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18. The compound according to Claim 1 represented by

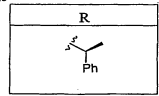
wherein R is

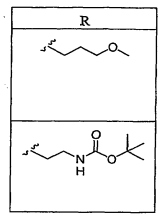
R



19. The compound according to Claim 1 represented by

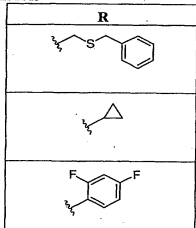
5 wherein R is





20. The compound according to Claim 1 represented by

5 wherein R is



21. The compound according to Claim 1 represented by

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wherein R is

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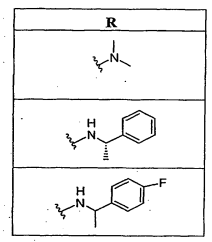
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or a pharmaceutically acceptable salt thereof.

25. A compound represented by

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or a pharmaceutically acceptable salt thereof.

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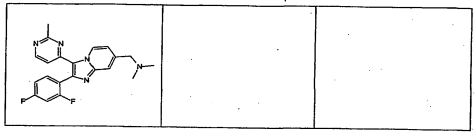
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	H ₂ N N N N N N N N N N N N N N N N N N N	NH ₂

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or a pharmaceutically acceptable salt thereof.

- 27. A pharmaceutical composition comprised of a compound in accordance with claim 1 in combination with a pharmaceutically acceptable carrier.
- 28. A method of treating a cytokine mediated disease in a mammal, comprising:

administering to a mammalian patient in need of such treatment a compound as described in claim 1 in an amount which is effective to treat said cytokine mediated disease.

- 29. A method of treating inflammation in a mammalian patient in need of such treatment, comprising:
- administering to said patient an anti-inflammatory effective amount of a compound as described in claim 1.
- 30. A method in accordance with claim 28 wherein the cytokine mediated disease is rheumatoid arthritis, osteoarthritis, endotoxemia, toxic shock syndrome, inflammatory bowel disease, tuberculosis, atherosclerosis, muscle degeneration, cachexia, psoriatic arthritis, Reiter's syndrome, rheumatoid arthritis, gout, traumatic arthritis, rubella arthritis or acute synovitis.
- 31. A method in accordance with claim 28 wherein the cytokine mediated disease is rheumatoid arthritis, rheumatoid spondylitis, osteoarthritis, gouty arthritis, sepsis, septic shock, endotoxic shock, gram negative sepsis, toxic shock syndrome, adult respiratory distress syndrome, cerebral malaria, chronic pulmonary inflammatory disease, silicosis, pulmonary sarcosis, bone resorption diseases,

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reperfusion injury, graft v. host rejection, allograft rejection, fever, myalgia due to infection, cachexia secondary to infection or malignancy, cachexia secondary to acquired immune deficiency syndrome (AIDS), AIDS related complex (ARC), keloid formation, scar tissue formation, Crohn's disease, ulcerative colitis or pyresis.

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- 32. A method of treating osteoporosis in a mammalian patient in need of such treatment, comprising administering to said patient an amount of a compound as described in claim 1 effective to treat osteoporosis.
- 10 33. A method of treating bone resorption in a mammalian patient in need of such treatment, comprising administering to said patient an amount of a compound as described in claim 1 effective to treat bone resorption.
 - 34. A method of treating Crohn's disease in a mammalian patient in need of such treatment comprising administering to said patient an amount of a compound as described in claim 1 effective to treat Crohn's disease.
 - 35. A method for the treatment or prevention of protozoal diseases comprising administering to a host in need of such treatment a therapeutically or prophylactically effective amount of a compound of Claim 1.
 - 36. A method for the treatment or prevention of coccidiosis in poultry comprising administering to the poultry a therapeutically or prophylactically effective amount of a compound of Claim 1.

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- 37. An antiprotoazoal composition comprising a compound of Claim 1 and an inert carrier.
- 38. A composition for the treatment or prevention of coccidiosis in poultry comprising a therapeutically or prophylactically effective amount of a compound of Claim 1 in poultry feedstuff.
 - 39. The composition of Claim 38, further comprising a second anticoccidial agent.

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- 40. A composition of Claim 39 wherein said second anticoccidial agent is selected from amprolium, ethopabate, clopidol, meticlorpindol, decoquinate, dinitolmide, halofuginone, lasalocid, maduramicin, monensin, narasin, nicarbazin, chlortetracycline, oxytetracycline, robenidine, salinomycin, semduramicin, and diclazuril.
- 41. A composition of Claim 39 wherein said second anticoccidial agent is selected from the group consisting of amprolium, ethopabate, lasalocid, monensin, salinomycin, and diclazuril.

INTERNATIONAL SEARCH REPORT

International application No.

PCT/US02/1950

		FC170302/1930/		
A. CLASSIFICATION OF SUBJECT MATTER IPC(7) : C07D 403/04, 417/04, 471/04; A61K 31/429, 31/506, 31/4355, 31/519, 31/53; A61P 19/02, 29/00 US CL : 544/331, 281, 184; 514/ 275, 258, 243 According to International Patent Classification (IPC) or to both national classification and IPC				
B. FIELDS SEARCHED				
Minimum documentation searched (classification system followed by classification symbols) U.S.: 544/331, 281, 184; 514/ 275, 258, 243				
Documentation searched other than minimum documentation to the extent that such documents are included in the fields searched				
Electronic data base consulted during the international search (name of data base and, where practicable, search terms used) CAS ONLINE, EAST				
C. DOCUMENTS CONSIDERED TO BE RELEVANT				
Category *	Citation of document, with indication, where a	propriate of the relevant passages	Relevant to claim No.	
	X WO 01/34605 A1 (ORTHO-MCNEIL PHARMACEUTICAL, INC.) 17 May 2001 (17.05.2001), see entire document especially pages 5-6, formula I and pages 9-11, compounds 9, 19, 20, and 21.			
			1, 3, 4	
		F-3	L	
Further	documents are listed in the continuation of Box C.	See patent family annex.		
"A" document defining the general state of the art which is not considered to be princip of particular relevance		date and not in conflict with the appli- principle or theory underlying the inv	cation but cited to understand the ention	
"B" earlier ap	plication or patent published on or after the international filing date	"X" document of particular relevance; the considered novel or cannot be considered novel or cann		
establish (specified)	•	considered to involve an inventive ste combined with one or more other suc	document of particular relevance; the claimed invention cannot be considered to involve an inventive step when the document is continued with one or more other such documents, such combination	
"O" document referring to an oral disclosure, use, exhibition or other means "D" document published prior to the international filing date but later than the		being obvious to a person skilled in the art		
"P" document published prior to the international filing date but later than the "& priority date claimed		"&" document member of the same patent family		
Date of the actual completion of the international search 12 September 2002 (12.09.2002)		Date of mailing of the international search report 25 SEP 2002		
Name and mailing address of the ISA/US Commissioner of Patents and Trademarks Box PCT Washington, D.C. 20231		Authorized officer Venkataraman Balasubramanian Telephone No. (703)308-1235		

Form PCT/ISA/210 (second sheet) (July 1998)